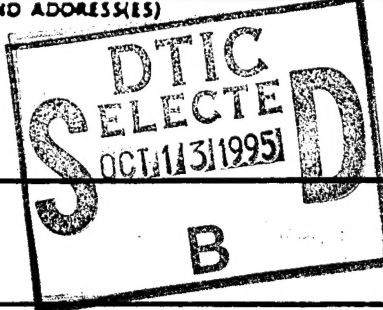


REPORT DOCUMENTATION PAGE

Public reporting burden for this collection of information is estimated to average 1 hour per response, including reviewing existing data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any aspect of this collection of information, including suggestions for reducing this burden, to Washington Headquarters Service, Directorate for Information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Paperwork Reduction Project (8704-0188), Washington, DC 20503.

1. AGENCY USE ONLY (Leave blank)	2. REPORT DATE 8/1/95	3. REPORT TYPE AND DATES COVERED Annual Report, 9/1/94-8/31/95
4. TITLE AND SUBTITLE (U) Studies on High Pressure and Unsteady Flame Phenomena		5. FUNDING NUMBERS PE - 61102F PR - 2308 SA - BS G - F49620-95-1-0092
6. AUTHOR(S) C. K. Law		
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) Princeton University Department of Mechanical and Aerospace Engineering Princeton, NJ 08544		8. PERFORMING ORGANIZATION REPORT NUMBER
9. SPONSORING/MONITORING AGENCY NAME(S) AND ADDRESS(ES) AFOSR/NA 110 Duncan Avenue, Suite B115 Bolling AFB DC 20332-0001		10. SPONSORING/MONITORING AGENCY REPORT NUMBER
11. SUPPLEMENTARY NOTES		
12a. DISTRIBUTION/AVAILABILITY STATEMENT Approved for public release; distribution is unlimited		12b. DISTRIBUTION CODE



ABSTRACT (Maximum 200 words)

The present study aims to gain fundamental understanding on the structure and response of steady and unsteady laminar premixed and nonpremixed flames in reduced and elevated pressure environments, and relate these understanding to the practical issues of flame extinction and turbulent combustion. The investigation herein has been conducted via the use of laser-based diagnostics, computational simulation of the flame structure with detailed chemistry and transport, and asymptotic analysis with reduced kinetic mechanisms. During the reporting period progress has been made in the following projects: (1) An analytical and experimental study of unsteady diffusion flames. (2) A computational and experimental study of the effects of thermophoresis on seeding particles in LDV measurements of flames. (3) A re-examination of the accuracy of the counterflow flame technique for the determination of laminar flame speeds. (4) Review articles written covering research performed under AFOSR support in recent years. A total of six reprints are appended.

DTIC QUALITY INSPECTED 5

14. SUBJECT TERMS Flames, high-pressure combustion, unsteady combustion, reduced mechanisms, extinction			15. NUMBER OF PAGES 107
			16. PRICE CODE
17. SECURITY CLASSIFICATION OF REPORT Unclassified	18. SECURITY CLASSIFICATION OF THIS PAGE Unclassified	19. SECURITY CLASSIFICATION OF ABSTRACT Unclassified	20. LIMITATION OF ABSTRACT UL

19951011 159

STUDIES ON HIGH PRESSURE AND UNSTEADY FLAME PHENOMENA

(AFOSR Grant No. F49620-95-1-0092)

Principal Investigator: Chung K. Law

Princeton University
Princeton, NJ 08544

SUMMARY/OVERVIEW

The objective of the present program is to study the structure and response of steady and unsteady laminar premixed and nonpremixed flames in reduced and elevated pressure environments through (a) non-intrusive experimentation, (b) computational simulation using detailed flame and kinetic codes, and (c) asymptotic analysis with reduced kinetic mechanisms. During the reporting period progress has been made in the following projects: (1) An analytical and experimental study of unsteady diffusion flames. (2) A computational and experimental study of the effects of thermophoresis on seeding particles in LDV measurements of flames. (3) A re-examination of the accuracy of the counterflow flame technique for the determination of laminar flame speeds. (4) Review articles written covering research performed under AFOSR support in recent years.

TECHNICAL DISCUSSIONS

1. Studies on Unsteady Diffusion Flames

An important influence on the flame behavior which so far has not been adequately addressed is the effect of unsteadiness of the environment on the flame behavior. This issue is of particular relevance to the modeling of turbulent flames through the concept of laminar flamelets. These flamelets are subject to fluctuating flows with various intensities of straining, and it is reasonable to expect that the flame would respond differently in an oscillating strained flow field than in a steady strained flow field.

During the reporting period we have first extended a previous asymptotic analysis on counterflow diffusion flames subject to *small-amplitude, sinusoidal* strain rate perturbations (Publication No. 1) to *arbitrary* strain rate as a function of time, with the assumptions of constant density and unity Lewis number. When specialized to the case of an impulsively-applied strain rate, the characteristic relaxation time as well as the extinction delay time are derived. For the sinusoidal strain rate, the reaction sheet oscillation is found to be asymmetrical with respect to its initial location, protruding more toward the freestream side. Furthermore, the amplitude of the oscillation decreases with frequency, and its phase delay approaches $\pi/2$ in the high frequency limit, as observed in previous experimental and numerical studies. For a given amplitude of oscillation, the flame is more easily extinguished when the characteristic time of oscillation is sufficiently long. When this result is applied to the current understanding of turbulent flames, it suggests that the laminar flame sheet can be sustained at higher Reynolds numbers. This is because there exists a range of eddies which, while possessing a sufficiently large strain rate to extinguish the flame in the steady limit, do not have sufficiently long characteristic time to effect extinction. Thus it seems reasonable to suggest that the applicable range of the laminar flamelet regime may be wider than can be expected from quasi-steady considerations.

The above concept is schematically shown in Fig. 1, in which we have plotted a normalized Damkohler number of an eddy versus the characteristic eddy size. The solid line represents the variation based on steady state considerations such that extinction of the eddy is expected when it crosses the maximum extinction Damkohler number line. However, allowing for unsteadiness, the

dotted lines show that crossing of this limit either is delayed or may not occur at all. Results from the above theoretical study are reported in Publication No. 2.

Experimentally, we have constructed a counterflow burner with strain rate oscillations being applied by loud speakers. An important consideration here is the ability to independently vary the frequency and amplitude of the oscillation. Preliminary results seem to indicate that, for a given frequency, extinction occurs at a constant maximum strain rate regardless of the steady-state strain rate. This would imply that extinction is a quasi-steady process, with the reaction zone only affected by the instantaneous strain rate it experiences. Unsteadiness, however, does seem to have a second order effect in that the instantaneous extinction strain rate is found to increase with frequency, as shown in Fig. 2.

2. Thermophoretic Effects on Seeding Particles in LDV Measurements of Flames

In a recent experimental and computational study on the detailed dynamic, thermal, and chemical structure of adiabatic, laminar counterflow premixed flames, we noticed that while close quantitative agreement between the measured and computed results exists for the scalar structure of the flame, the LDV-measured axial velocity profile consistently lags the calculated values by substantial amounts in the preheat zone of the flame, as shown in Fig. 3. Order of magnitude estimates showed that such a lag could be due to the influence of thermophoresis on the LDV seeding particles in the high-temperature-gradient environment of the preheat zone. Indeed, when detailed calculations were performed for the motion of the seeding particles, under the influence of drag and thermophoresis and by using the computationally-determined flame structure to evaluate the various transport coefficients, the computed particle trajectory agreed well with the measured LDV velocity profile, as shown in Fig. 3.

Since temperature increases monotonically in the direction of the flow in an adiabatic *premixed* flame, the effect of thermophoresis can be readily visualized as in Fig. 3. However, for a counterflow *diffusion* flame, the temperature peaks in the flow field such that the thermophoretic force acts in opposite direction in the fuel and oxidizer sides of the flame. Furthermore, since the flow is uni-directional in crossing the flame, and the direction also switches when the flame moves across the stagnation surface, the net dynamic response of the LDV particles can be very rich. Figure 4 shows the evolution of the measured particle velocity profiles and computed gas velocity profiles for flames with the same calculated adiabatic flame temperature but different stoichiometric mixture fractions and hence locations relative to the stagnation surface. The richness and complexity with which the particle velocity can be modified by thermophoresis is quite evident.

There are several implications of this finding. First, thermophoresis appears to be a factor that needs to be estimated and possibly accounted for when measuring flame properties and responses using LDV and PIV, especially for thin flames in local flow field of low convective velocities. Its potential influence on measuring the velocity statistics in turbulent flames also needs to be examined. Furthermore, caution is also needed in the study of flame chemistry by directly extracting the local temperature profile from the local LDV-velocity profile, without considering thermophoresis.

The above work is reported in Publication Nos. 3 and 4.

3. Re-Examination of the Counterflow Technique in Laminar Flame Speed Determination

The accuracy of the laminar flame speed determination by using the counterflow twin flame technique has been computationally and experimentally examined in light of the recent understanding that linear extrapolation of the reference upstream velocity to zero strain rate would yield a value higher than that of the laminar flame speed, and that such an over-estimate can be reduced by using either lower strain rates and/or larger nozzle separation distances. A systematic evaluation of the above concept has been conducted and verified for the ultra lean hydrogen/air flames which have relatively large Karlovitz numbers, even for small strain rates, because of their very small laminar flame speeds. Consequently, the significantly higher values of the previous experimentally measured flame speeds, as compared to the independently calculated laminar flame speeds, can now be attributed to the use of nozzle separation distances which were not sufficiently large and/or strain rates which were not sufficiently small. Thus by using lower strain rates and larger nozzle separation distances the experimentally and computationally re-determined values of these ultra lean hydrogen/air flames agree well with the calculated laminar flame speeds (Fig. 5).

Dist		Special	
A-1			

The laminar flame speeds of methane/air and propane/air mixtures have also been experimentally re-determined over extensive ranges of the equivalence ratio and are found to be slightly lower than previously reported experimental values. Figure 6 shows the data and comparison for the propane/air flames.

This work is reported in Publication No. 5.

4. Review Articles

Two major review articles have been written on combustion phenomena whose understanding has been made possible through several long-term research programs including the present one. The first is on microgravity combustion (Publication No. 6) in which interpretation of the flame structure and aerodynamic response reached fruition through the present program. The second is on the role of chain mechanisms in combustion phenomena (Publication No. 6) in which chemical and aerodynamic effects on the flame structure and response are discussed from a unified viewpoint.

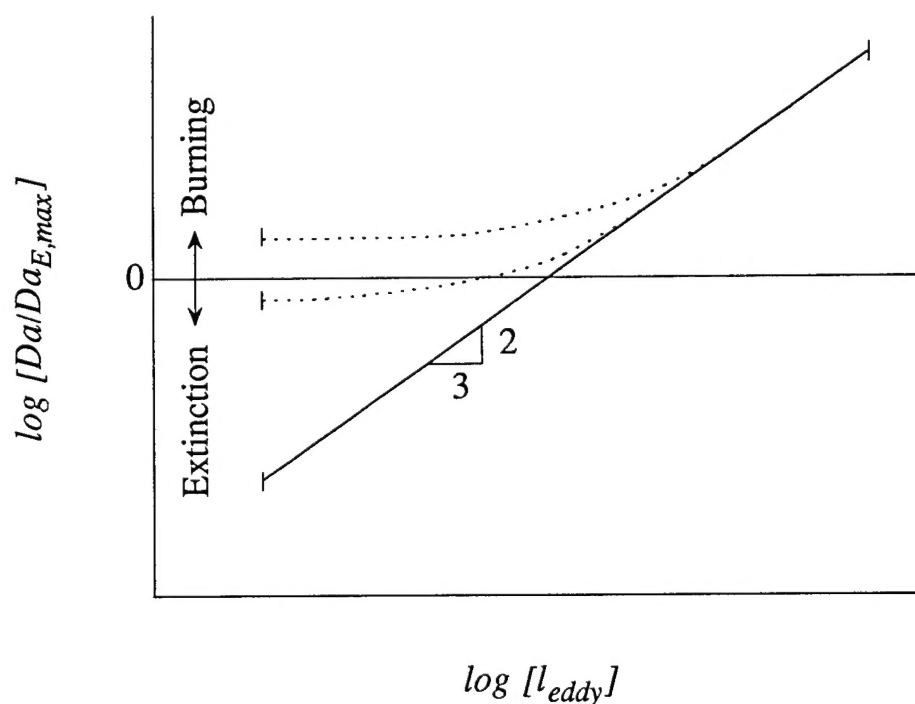


Figure 1

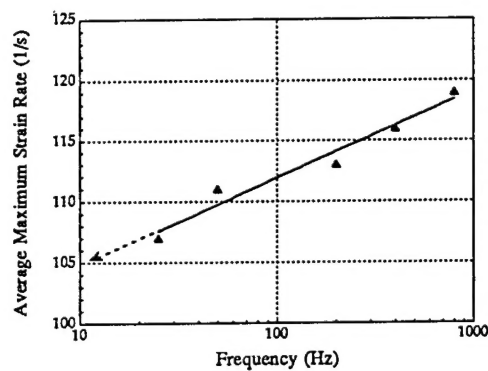


Figure 2

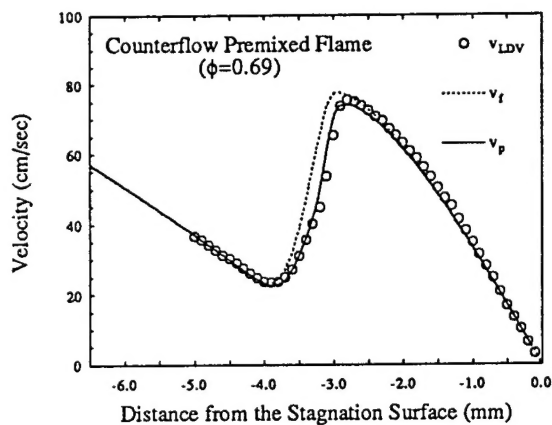


Figure 3

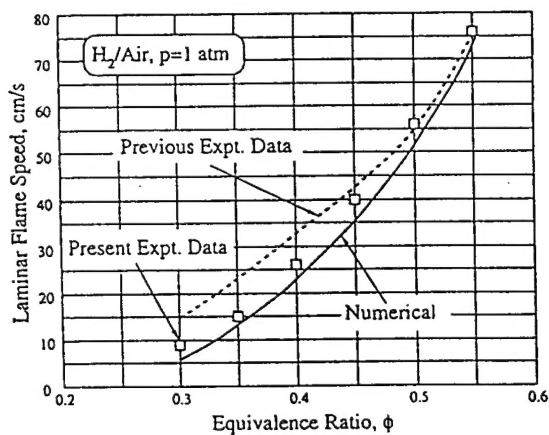


Figure 5

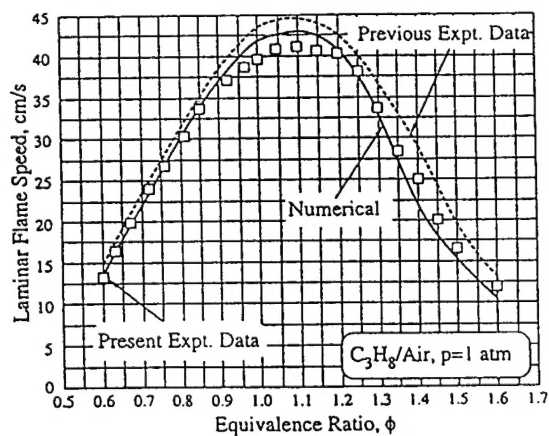


Figure 6

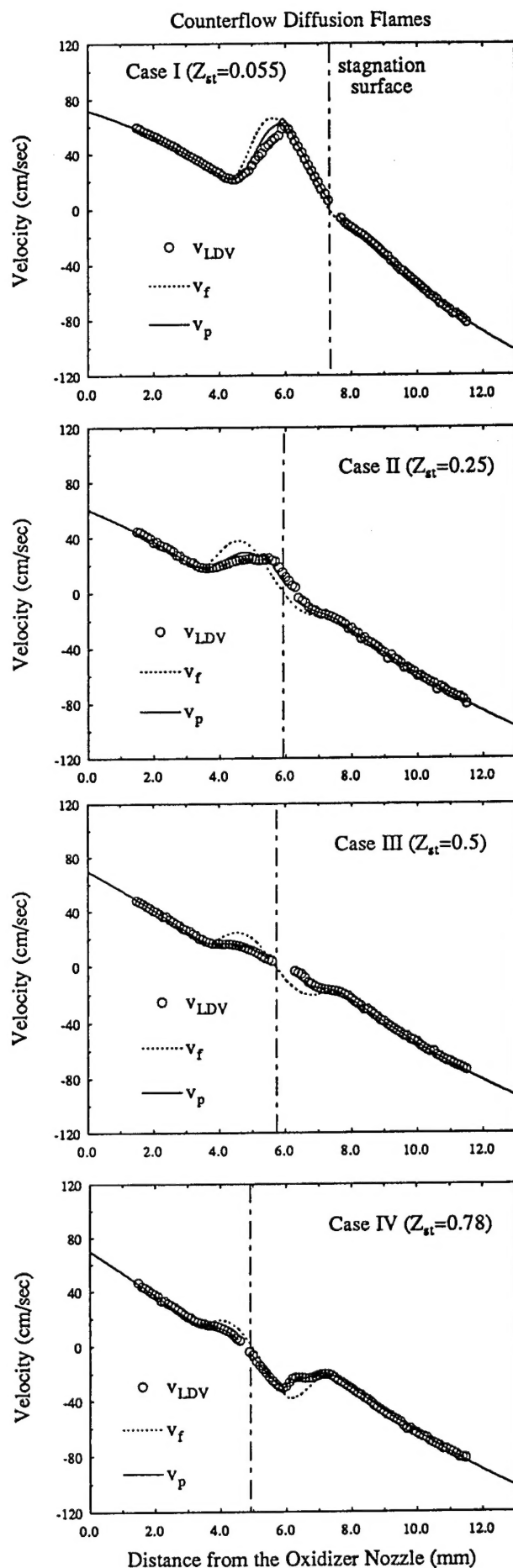


Figure 4

MAJOR PUBLICATIONS

1. "Response of counterflow diffusion flames to oscillating strain rates," by H. G. Im, C. K. Law, J. S. Kim and F. A. Williams, *Combustion and Flame*, Vol. 100, pp. 21-30 (1995). [Attached as Appendix A]
2. "Counterflow diffusion flames with unsteady strain rates," by H.G. Im, J.K. Bechtold and C.K. Law, *Combustion Science and Technology*, in press.
3. "Thermophoretic effects of seeding particles in LDV measurements of flames," by C.J. Sung, C.K. Law and R.L. Axelbaum, *Combustion Science and Technology*, Vol. 99, pp. 119-132 (1994). [Attached as Appendix B]
4. "Further studies on effects of thermophoresis on seeding particles in measurements of strained flames," by C.J. Sung, J.S. Kistler, M. Nishioka and C.K. Law, *Combustion and Flame*, in press.
5. "Further considerations on the determination of laminar flame speeds with the counterflow twin flame technique," by C.M. Vagelopoulos, F.N. Egolfopoulos and C.K. Law, *Twenty-Fifth Symposium (International) on Combustion*, pp. 1341-1347 (1994). [Attached as Appendix C]
6. "Opportunities and challenges of combustion in microgravity," by C.K. Law and G.M. Faeth, *Progress in Energy and Combustion Science*, Vol. 20, pp. 65-113 (1994). [Attached as Appendix D]
7. "The role of chain mechanisms in some fundamental combustion phenomena," by C.K. Law, submitted.
8. "Extinction of premixed methane-air flames with volumetric heat loss," by J.K. Bechtold and C.K. Law, *Combustion Science and Technology*, Vol. 100, pp. 371-383 (1994). [Attached as Appendix E]
9. "Effects of pressure and dilution on the extinction of counterflow nonpremixed hydrogen-air flames," by P. Papas, I. Glassman and C.K. Law, *Twenty-Fifth Symposium (International) on Combustion*, pp. 1333-1339 (1994). [Attached as Appendix F]

MEETING PREPRINTS AND PRESENTATIONS

1. "Counterflow diffusion flames with unsteady strain rates," by H.G. Im, J.K. Bechtold and C.K. Law, AIAA-95-0128, 33rd Aerospace Space Sciences Meeting, Reno, NV, Jan. 9-12, 1995.

PERSONNEL

1. H.G. Im (Doctoral student, graduated and assumed a post-doctoral position at the Stanford/Ames Center for Turbulence Research in September, 1995)
2. J.K. Bechtold (Research staff, assumed an assistant professorship in mathematics at the New Jersey Institute of Technology in September, 1995)
3. C.J. Sung (Doctoral student, graduated and assumed a research staff position at Princeton University in September, 1995)
4. M. Nishioka (Visiting Fellow from Nagoya University, Japan)
5. J.S. Kistler (MS student, graduated in June, 1995)

HONORS AND AWARDS

1. 1994 AIAA Propellants and Combustion Award
2. Appointment to the Robert H. Goddard Professorship at Princeton University

Technology Transitions

1. Performer: C. K. Law; Princeton University;
(609) 258-5271.

Customer: Gas Research Institute; R. Serauskas;
(312) 399-8208.

Result: Accurate experimental determination of
laminar flame speeds of combustible
mixtures.

Application: Development and application of
GRI-Mech mechanism for methane
oxidation.

2. Performer: C. K. Law; Princeton University;
(609) 258-5271.

Customer: E. I. Dupont Company; P. Gelblum;
(302) 774-2249.

Result: Theory for determination of flammability
limits of combustible mixtures

Application: Assessment of explosion limits in chemical
processing.

Response of Counterflow Diffusion Flames to Oscillating Strain Rates

Appendix A

H. G. IM and C. K. LAW

*Department of Mechanical and Aerospace Engineering
Princeton University
Princeton, NJ 08544*

J. S. KIM and F. A. WILLIAMS

*Department of Applied Mechanics and Engineering Sciences
University of California, San Diego
La Jolla, CA 92093*

The response of counterflow flames to oscillating strain rates is analyzed by using large activation energy asymptotics, as a potential application to turbulent combustion and acoustic instability of rocket engines. The characteristic oscillation time of practical interest is found to be of the same order as the characteristic diffusion time of the flame, so that the flame structure consists of a quasi-steady reactive-diffusive layer embedded in the outer unsteady-diffusive-convective zone. A linear analysis is conducted by assuming that the amplitude of the strain rate oscillation is small relative to the mean strain rate. Results show that the flame response is controlled mainly by two effects: (a) the response of the convective mass flux into the reaction sheet, which is directly related to the flow-field variation applied at the boundary, and (b) the response of the reaction sheet to adjust the reduced residence time due to finite-rate chemistry. For flames near equilibrium, the former effect tends to be dominant, so that the response of the net heat release is in phase with the strain rate oscillation. For flames near extinction, however, the finite-rate chemistry effect overtakes the fluid-dynamic effect such that increasing strain rate leads to a reduction of the reactivity of the flame during the oscillatory cycle. As such, the net heat release response of the near-extinction flame becomes out of phase with the strain rate oscillation in the sense of the Rayleigh's criterion. Results of the present study suggest the possibility that the unsteady characteristics of the near-extinction diffusion flame can be significantly different from those in the Burke-Schumann limit.

INTRODUCTION

Effects of unsteadiness on the laminar flame characteristics have recently received attention in an attempt to understand unsteady phenomena occurring in turbulent combustion [1-4]. Since a turbulent flow consists of eddies with a wide spectrum of length and time scales, it is expected that if a turbulent Reynolds number is sufficiently large, there exists a range of eddy sizes in which the characteristic eddy turnover time becomes comparable with the longest characteristic time, i.e., the diffusion time, of the laminar flamelet. Typically, the eddies with sizes near the Gibson scale of turbulent premixed flames [5], or eddies that can cause extinction of diffusion flames [6] fall into this

category. Under these situations, the unsteadiness begins to show its effect on the characteristics of laminar flamelets. As a first step to address such an effect, it is appropriate to analyze the behavior of laminar flames modified by unsteadiness arising from a small amplitude, monochromatic oscillation of the strain rate with respect to its mean value, recognizing, however, that the inherent randomness involved in turbulence may significantly complicate the direct application of the present result to the laminar flamelet regime of turbulent combustion.

The response of diffusion flames to a monochromatic oscillatory strain rate is also relevant to acoustic instability phenomena occurring in liquid-propellant rocket engines. The condition under which acoustic instability occurs is well understood through the Rayleigh's criterion, which states that acoustic amplification occurs if, on the average, heat is added in

Presented at the Twenty-Fifth Symposium (International) on Combustion, Irvine, California, 31 July-5 August 1994.

COMBUSTION AND FLAME 100: 21-30 (1995)
Copyright © 1995 by The Combustion Institute
Published by Elsevier Science Inc.

0010-2180/95/\$9.50
SSDI 0010-2180(94)00059-2

phase with the pressure increase during the oscillation. Since acoustic pressure oscillations always accompany the corresponding velocity oscillations, acoustic waves normally influence flames through both pressure and velocity effects. The response of strained diffusion flames to acoustic pressure oscillations has recently been studied by Kim and Williams [6]. The present investigation then aims to complement this previous study by providing the corresponding velocity responses. In particular, we examine the flame response coupled with finite-rate chemistry, so that the present study is distinguished from some previous works [7, 8] in which the flame is only treated in the Burke-Schumann limit.

In this study we analyze the laminar counterflow diffusion flame subject to an oscillating strain rate with small amplitude. We use asymptotic analysis employing a one-step irreversible Arrhenius reaction with a large activation energy, which corresponds to a large Zel'dovich number β . For such a system, chemical reaction is confined to an asymptotically thin layer of $O(\beta^{-1})$ compared with the diffusive transport zone, and thus the characteristic time for reaction is much shorter, by $O(\beta^{-2})$, than the characteristic diffusion time. If we focus attention on flames near extinction, the characteristic diffusion time of the laminar diffusion flame can be estimated by the extinction strain rate of the stoichiometric hydrocarbon/air flame in the range of 10^{-3} s, which turns out to be of the same order as the characteristic unsteady time often encountered in practical problems such as acoustic instability in rocket engines or turbulent reacting flows [6, 9]. Therefore, in the present analysis the time scale of the unsteadiness is chosen to be comparable to that of diffusive transport, such that the outer diffusive-convective layer is modified to include the unsteadiness caused by the oscillating strain rate, while the inner reactive-diffusive layer remains quasi-steady. The effect of finite-rate chemistry will then influence the unsteady flame response through instantaneous matching conditions. We particularly wish to predict the response of the reaction sheet and the burning rate to time-varying strain rates in terms of their magnitude and phase. It will be demonstrated that such re-

sponses become very sensitive to flow unsteadiness as the mean flame approaches the extinction condition.

ANALYSIS

Formulation

As illustrated in Fig. 1, we consider an unsteady, axisymmetric counterflow diffusion flame in the vicinity of the axis of symmetry. In terms of the mean radial velocity gradient in the external oxidizer stream a_∞ , the nondimensional similarity variable for the axial coordinate and the nondimensional time are, respectively, defined as

$$\eta \equiv \left(\frac{2a_\infty}{\nu_\infty} \right)^{1/2} \int_{z_s}^z \frac{\rho(z', \tau)}{\rho_\infty} dz', \quad \tau \equiv 2a_\infty t, \quad (1)$$

where t is the time, ρ the density, ν the kinematic viscosity, z_s the location of the stagnation plane, and the subscripts " ∞ " and " $-\infty$," respectively, denote conditions at the external oxidizer and fuel streams.

Since the radial velocity component u is proportional to the radius r , u is related to the mean radial velocity gradient in the external oxidizer stream a_∞ and to the nondimensional unsteady stream functions $f(\eta, \tau)$ by $u = a_\infty r f_\eta$, where $()_\eta \equiv \partial/\partial\eta$, etc. The oscillation of the

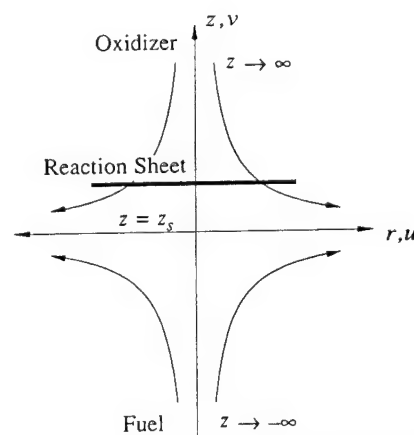


Fig. 1. Schematic of the counterflow diffusion flame.

strain rate is then imposed as the boundary condition for the radial velocity at the external oxidizer stream given by

$$f_{\eta,\infty} = f_{\eta}(\infty, \tau) = 1 + \alpha e^{i\omega\tau}, \quad (2)$$

where α is the relative amplitude of strain rate fluctuation to the mean strain rate, and ω the dimensional frequency divided by $2a_x$. The radial velocity in the fuel stream also oscillates in a manner that the momentum balance is maintained across the viscous boundary layer.

Assuming unity Prandtl number, constant $\rho\mu$, and constant average molecular weight, the conservation equations for radial momentum, species and temperature can then be written as

$$f_{\eta\eta\eta} + ff_{\eta\eta} + \frac{1}{2}[(1 + q\theta)F(\tau) - f_{\eta}^2 - 2f_{\eta\tau}] = 0, \quad (3)$$

$$\frac{1}{Le_i} Y_{i\eta\eta} + fY_{i\eta} - Y_{\tau} = w \quad \text{for} \quad i = F, O, \quad (4)$$

$$\theta_{\eta\eta} + f\theta_{\eta} - \theta_{\tau} = -w, \quad (5)$$

where

$$\theta = c_p(T - T_x)/Q \equiv (T - T_x)/(qT_x) \quad (6)$$

is the nondimensional temperature, c_p the specific heat, Q the heat release per unit mass of fuel, $q \equiv Q/c_p T_x$ the nondimensional heat release, Y_F the fuel mass fraction, Y_O the oxidizer mass fraction scaled by the stoichiometric oxidizer-to-fuel mass ratio σ , and $F(\tau) = (f_{\eta,\infty})^2 + 2(f_{\eta,\infty})_{\tau}$ is a function determined by the imposed boundary condition in Eq. 2. The chemical source term in Eqs. 4 and 5 is

$$w = Da Y_F Y_O \exp(-E/RT), \quad (7)$$

where E is the activation energy, $Da \equiv B\sigma/2a_x$ the Damköhler number, and B the frequency factor. The reaction orders with respect to both fuel and oxidizer are taken to be unity.

The above system of equations is subject to the boundary conditions

$$\begin{aligned} f_{\eta}^2 + 2f_{\eta\tau} &\rightarrow F(\tau), \quad Y_F \rightarrow 0, \quad Y_O \rightarrow Y_{O,\infty}, \\ \theta &\rightarrow 0, \quad \text{as } \eta \rightarrow \infty, \\ f_{\eta}^2 + 2f_{\eta\tau} &\rightarrow (1 + q\theta_{-\infty})F(\tau), \quad Y_F \rightarrow Y_{F,-\infty}, \\ Y_O &\rightarrow 0, \quad \theta \rightarrow \theta_{-\infty}, \quad \text{as } \eta \rightarrow -\infty. \end{aligned} \quad (8)$$

In addition, $f = 0$ at $\eta = 0$ because the origin of the η coordinate is placed at the stagnation plane.

Analysis of the Outer, Nonreactive Field

For a system with a large activation energy, reaction is confined to an asymptotically thin reaction sheet, and the rest of the transport zone is nonreactive to all algebraic orders of the small parameter β^{-1} . The flame response in this outer, nonreactive zone can be studied by conducting a linear analysis for a small oscillation amplitude, $\alpha \ll 1$. Given the oscillating boundary condition in Eq. 2, the reaction-sheet location η_f is expected to respond as

$$\eta_f = \bar{\eta}_f + \alpha e^{i\omega\tau} \tilde{\eta}_f + O(\alpha^2), \quad (9)$$

where $\tilde{\eta}_f$ is the complex amplitude of the reaction-sheet oscillation of $O(1)$. Any dependent variable \mathcal{F} is also expanded in a form [6]

$$\begin{aligned} \mathcal{F}(\eta; \eta_f) &= \bar{\mathcal{F}}(\eta; \bar{\eta}_f) + \alpha e^{i\omega\tau} \\ &\times [\tilde{\mathcal{F}}(\eta; \bar{\eta}_f) + \tilde{\eta}_f \hat{\mathcal{F}}(\eta; \bar{\eta}_f)] + O(\alpha^2), \end{aligned} \quad (10)$$

where $\hat{\mathcal{F}}(\eta; \eta_f) \equiv \partial \bar{\mathcal{F}} / \partial \eta_f$. Here the term involving $\tilde{\mathcal{F}}$ represents the direct influence of the oscillatory strain rate, while the term involving $\tilde{\eta}_f \hat{\mathcal{F}}$ represents the indirect effect that arises through oscillation of the reaction sheet. Substituting the above expansions into Eqs. 3–5, and collecting terms of the same order in α , we obtain the governing equations for the mean and oscillatory states.

At the leading order in α , the conservation equations for the mean state become

$$\bar{f}_{\eta\eta\eta} + \bar{f}\bar{f}_{\eta\eta} + \frac{1}{2}(1 + q\bar{\theta} - \bar{f}_{\eta}^2) = 0, \quad (11)$$

$$\frac{1}{Le_i}\bar{Y}_{i\eta\eta} + \bar{f}\bar{Y}_{i\eta} = 0, \quad (12)$$

$$\bar{\theta}_{\eta\eta} + \bar{f}\bar{\theta}_{\eta} = 0, \quad (13)$$

which are subject to the boundary conditions

$$\begin{aligned} \bar{f}_{\eta} &= 1, \quad \bar{Y}_F = 0, \quad \bar{Y}_O = Y_{O,\infty}, \\ \bar{\theta} &= 0, \quad \text{as } \eta \rightarrow \infty, \\ \bar{f}_{\eta} &= (1 + q\theta_{-\infty})^{1/2}, \quad \bar{Y}_F = Y_{F,-\infty}, \quad \bar{Y}_O = 0, \\ \bar{\theta} &= \theta_{-\infty}, \quad \text{as } \eta \rightarrow -\infty, \end{aligned} \quad (14)$$

and the jump conditions across the reaction sheet

$$\begin{aligned} \left[\bar{\theta}_{\eta} + \frac{1}{Le_i}\bar{Y}_{i\eta} \right]_{\bar{\eta}_f}^{\bar{\eta}_f^+} &= 0, \quad \text{for } i = F, O, \\ \bar{Y}_F|_{\bar{\eta}_f^-} &= \bar{Y}_F|_{\bar{\eta}_f^+} = \bar{Y}_F|_{\bar{\eta}_f}, \quad \bar{Y}_O|_{\bar{\eta}_f^-} = \bar{Y}_O|_{\bar{\eta}_f^+} = 0, \\ \bar{\theta}|_{\bar{\eta}_f^-} &= \bar{\theta}|_{\bar{\eta}_f^+} = \bar{\theta}|_{\bar{\eta}_f}. \end{aligned} \quad (15)$$

At the next order in α , we have the governing equations for the unsteady fluctuations as

$$\begin{aligned} \tilde{f}_{\eta\eta\eta} + \tilde{f}\tilde{f}_{\eta\eta} + \tilde{f}\tilde{f}_{\eta\eta} + \left[(1 + i\omega)(1 + q\bar{\theta}) \right. \\ \left. + \frac{1}{2}q\bar{\theta} - \bar{f}_{\eta}\tilde{f}_{\eta} \right] - i\omega(\tilde{f}_{\eta} + \tilde{\eta}_f\tilde{f}_{\eta}) &= 0, \end{aligned} \quad (16)$$

$$\frac{1}{Le_i}\tilde{Y}_{i\eta\eta} + \tilde{f}\tilde{Y}_{i\eta} + \tilde{f}\tilde{Y}_{i\eta} - i\omega(\tilde{Y}_i + \tilde{\eta}_f\tilde{Y}_i) = 0, \quad (17)$$

$$\tilde{\theta}_{\eta\eta} + \tilde{f}\tilde{\theta}_{\eta} + \tilde{f}\tilde{\theta}_{\eta} - i\omega(\tilde{\theta} + \tilde{\eta}_f\tilde{\theta}) = 0, \quad (18)$$

with the boundary conditions

$$\begin{aligned} \tilde{f}_{\eta} &= 1, \quad \tilde{Y}_i = \tilde{\theta} = 0, \quad \text{as } \eta \rightarrow \infty, \\ \tilde{f}_{\eta} &= \frac{(1 + i\omega)(1 + q\theta_{-\infty})}{(1 + q\theta_{-\infty})^{1/2} + i\omega}, \quad \tilde{Y}_i = \tilde{\theta} = 0, \\ \text{as } \eta &\rightarrow -\infty. \end{aligned} \quad (19)$$

The jump conditions are identical to those for the steady system, Eq. 15, obtained by replacing $\bar{\theta}$, \bar{Y}_i by $\tilde{\theta}$ and \tilde{Y}_i .

To close the system of Eqs. 16–18, an additional condition to determine the eigenvalue $\tilde{\eta}_f$ for a given $\bar{\eta}_f$ is provided by analyzing the reactive-diffusive layer.

Analysis of the Inner, Reactive-Diffusive Zone

Within the thin inner layer, reaction and diffusion are balanced in a quasi-steady manner. In addition, for most practical diffusion flames employing hydrocarbon or hydrogen as fuel against air, the stoichiometric mixture fraction is so small that the reaction sheet is situated far into the oxidizer stream. Under this condition, we adopt Liñán's premixed flame regime analysis [10], in which the abundant species, namely the fuel, is assumed to leak through the reaction sheet by an $O(1)$ amount.

In the inner reactive layer, the appropriate stretched variables are defined by

$$\begin{aligned} \theta_{in} &= \theta|_{\eta_f} - \beta^{-1}(\vartheta + m\xi), \\ Y_{O,in} &= \beta^{-1}Le_O\phi, \quad Y_{F,in} = Y_F|_{\eta_f} + O(\beta^{-1}), \end{aligned} \quad (20)$$

and

$$\xi = \beta Le_O^{-1} Y_{O\eta}|_{\eta_f^+} (\eta_f - \eta) + (\vartheta - \phi)_{\infty}/m, \quad (21)$$

where the small parameter of expansion is the reciprocal of the Zel'dovich number

$$\beta \equiv qE / [RT_{\infty}(1 + q\theta|_{\eta_f})^2]. \quad (22)$$

The parameter m in Eqs. 20 and 21 is

$$m = Le_O(\theta_{\eta}|_{\eta_f^-}) / (Y_{O\eta}|_{\eta_f^+}), \quad (23)$$

which is the fractional amount of the total heat release lost to the fuel side.

Substituting the inner variables into the governing equations, we obtain the canonical

equation and boundary conditions for the inner structure as

$$\phi_{\xi\xi} = \Lambda \phi \exp[-(\phi + m\xi)],$$

$$\phi_{\xi} \rightarrow 0 \text{ as } \xi \rightarrow \infty, \phi_{\xi} \rightarrow -1 \text{ as } \xi \rightarrow -\infty, \quad (24)$$

where Λ is the reduced Damköhler number defined as

$$\Lambda = \text{Da} \frac{\text{Le}_O^3 Y_F|_{\eta_f}}{\beta^2 (Y_{O\eta}|_{\eta_f})^2} \exp\left[\frac{-\beta}{1 + q\theta|_{\eta_f}}\right], \quad (25)$$

which is an eigenvalue of the problem. Equation 24 has been solved numerically, and the eigenvalue Λ is found to be a function of the parameter m , which has been numerically fitted [10] as

$$\Lambda \approx (1 - 1.344m + 0.6307m^2)/2. \quad (26)$$

Finally, the additional eigenvalue $\tilde{\eta}_f$ can be determined by equating the unsteady parts of Eqs. 25 and 26, which is equivalent to the requirement that the Damköhler number Da does not vary with time [6]. This leads to

$$\bar{\Delta} + \tilde{\eta}_f \bar{\Delta} = 0, \quad (27)$$

where

$$\bar{\Delta} = \frac{\bar{m}}{\bar{\Lambda}} \frac{d\bar{\Lambda}}{d\bar{m}} \left[\frac{\bar{\theta}_{\eta}|_{\bar{\eta}_f}}{\bar{\theta}_{\eta}|_{\bar{\eta}_f}} - \frac{\bar{Y}_{O\eta}|_{\bar{\eta}_f}}{\bar{Y}_{O\eta}|_{\bar{\eta}_f}} \right] + \frac{2\bar{Y}_{O\eta}|_{\bar{\eta}_f}}{\bar{Y}_{O\eta}|_{\bar{\eta}_f}} - \frac{\bar{Y}_F|_{\bar{\eta}_f}}{\bar{Y}_F|_{\bar{\eta}_f}} - \frac{4q\bar{\theta}|_{\bar{\eta}_f}}{1 + q\bar{\theta}|_{\bar{\eta}_f}} - \bar{\beta}\bar{\theta}|_{\bar{\eta}_f},$$

$$\bar{\Delta} = \frac{\bar{m}}{\bar{\Lambda}} \frac{d\bar{\Lambda}}{d\bar{m}} \left[\frac{\bar{\theta}_{\eta}|_{\bar{\eta}_f}}{\bar{\theta}_{\eta}|_{\bar{\eta}_f}} - \frac{\bar{Y}_{O\eta}|_{\bar{\eta}_f}}{\bar{Y}_{O\eta}|_{\bar{\eta}_f}} \right] + \frac{2\bar{Y}_{O\eta}|_{\bar{\eta}_f}}{\bar{Y}_{O\eta}|_{\bar{\eta}_f}} - \frac{\bar{Y}_F|_{\bar{\eta}_f}}{\bar{Y}_F|_{\bar{\eta}_f}} - \frac{4q\bar{\theta}|_{\bar{\eta}_f}}{1 + q\bar{\theta}|_{\bar{\eta}_f}} - \bar{\beta}\bar{\theta}|_{\bar{\eta}_f}. \quad (28)$$

Here “ $\bar{\cdot}$ ” denotes the total derivative of a quantity with respect to η_f at $\bar{\eta}_f$, i.e.,

$$\bar{\mathcal{F}}|_{\bar{\eta}_f} = \frac{d\bar{\mathcal{F}}}{d\eta_f}\bigg|_{\bar{\eta}_f} = \bar{\mathcal{F}}_{\eta}(\bar{\eta}_f; \bar{\eta}_f) + \hat{\mathcal{F}}(\bar{\eta}_f; \bar{\eta}_f), \quad (29)$$

so that $\bar{\Delta}$ represents the fractional variation of Da due to the fluctuation of the unsteady field and $\bar{\Delta}$ due to the unsteady response of the reaction sheet.

In summary, the calculation procedure is the following: first we select a mean reaction-sheet location $\bar{\eta}_f$ and solve the steady system Eqs. 11–15. Then using the results of the inner structure analysis, Eqs. 22, 23, 25, and 26, we find the corresponding Damköhler number, Da . Next, we proceed with the unsteady system, Eqs. 16–19, plus the additional constraint, Eq. 27, to determine the oscillatory field and $\tilde{\eta}_f$ simultaneously. By repeating the same procedure for different values of $\bar{\eta}_f$, the flame structure as well as its response to the oscillating strain rate can be calculated as a function of the system Damköhler number.

RESULTS AND DISCUSSIONS

The parameter values used in the present calculations are $q = 50$, $E/RT_{\infty} = 50$, $\theta_{-\infty} = 0$, $Y_{F,-\infty} = 1.0$, and $Y_{O,\infty} = 0.1$. The Lewis numbers for both fuel and oxidizer are taken to be unity, except for the last case in which we compare the results for various Le_F .

Figure 2 shows a typical result of the steady reaction-sheet response as a function of Da . It is seen that for large Da , the steady reaction-sheet is close to the Burke-Schumann limit, so that neither reactant leaks through the reaction sheet by $O(1)$ amount. As the strain rate

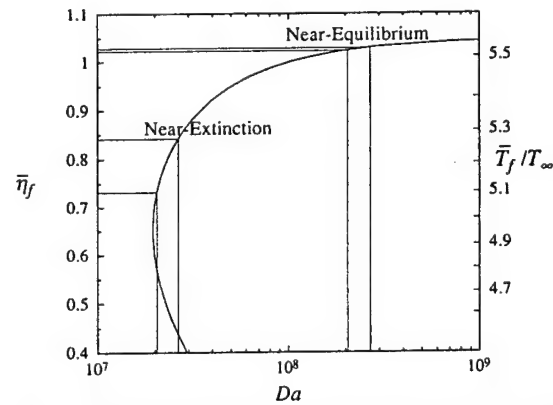


Fig. 2. Steady reaction-sheet location as a function of the strain rate, showing the sensitivity of the flame response near extinction.

increases, the reaction sheet moves toward the stagnation plane and finally exhibits a turning point that corresponds to the static extinction point. In the present calculation, the static extinction occurs at $\bar{\eta}_f = 0.653$, $\bar{T}_f/T_\infty = 4.953$ and $Da = 1.952 \times 10^7$. The well-known steady S-curve behavior in Fig. 2 illustrates that, for the same amount of the fractional variation of the Damköhler number $\delta Da/Da$, shown as vertical bars, the near-extinction flame yields much larger reaction-sheet displacement and the flame temperature variation, as marked by horizontal bars, than the near-equilibrium flame. Figure 2 therefore provides useful insight on the sensitivity of the flame response to the oscillatory strain rate, even the reaction sheet does not exactly follow the steady S-curve when oscillatory strain is imposed.

Figure 3 shows the behavior of the real part of the reaction-sheet fluctuation, $Re(\tilde{\eta}_f)$, as a function of the normalized Damköhler number for various ω , where Da_E is the static extinction Damköhler number. It is first noted that, for all ω , $Re(\tilde{\eta}_f)$ is negative throughout the entire range of the Damköhler number, indicating that the reaction sheet moves toward the stagnation plane when the strain rate is increased with time. Specifically, during a period of strain rate oscillation, an increase in strain rate results in reduced residence time in the reaction zone, thereby reducing the reaction rate. In order to compensate for the re-

duced reactivity, the reaction sheet migrates toward the stagnation plane such that the local convective flux normal to the reaction sheet is reduced. Therefore, the negative values of $Re(\tilde{\eta}_f)$ indicates the reduced/increased reactivity when the strain rate is increased/decreased during the oscillatory cycle.

As expected from the steady flame response in Fig. 2, Fig. 3 shows that the reaction-sheet fluctuation is greatly amplified as the steady flame nears extinction. The reaction-sheet response, however, does not grow indefinitely at the extinction point because the increase of $\tilde{\eta}_f$ is fed into the source terms in the unsteady equations 16–18, resulting in a smaller value of \tilde{A} . Therefore, both \tilde{A} and \tilde{B} in Eq. 27 vanish at extinction, and a finite value of $\tilde{\eta}_f$ is achieved, provided that ω is not identically zero [6]. We also remark that, although not shown here, the absolute magnitude of $\tilde{\eta}_f$ exhibits behavior similar to the real parts plotted in Fig. 3. As for the dependence on frequency, the sensitive behavior of the reaction sheet near extinction is seen to be more prominent for lower frequencies, because the flame has sufficient time to respond to the external fluctuation as its frequency becomes smaller.

We next examine the response of the heat release rate, or equivalently reactant consumption rate, to the unsteady strain rate. The normalized net heat release rate can be defined by the instantaneous reactant gradient at the reaction sheet, namely

$$h_{net} = Y_{O\eta}|_{\eta_f^*} \quad (30)$$

Then the oscillating part of the heat release rate scaled by α can be decomposed into

$$\tilde{h}_{net} = \tilde{h}_c + \tilde{h}_r = \tilde{Y}_{O\eta}|_{\tilde{\eta}_f^*} + \tilde{\eta}_f(\tilde{Y}_{O\eta\eta} + \hat{Y}_{O\eta})_{\tilde{\eta}_f^*}, \quad (31)$$

where \tilde{h}_c represents the contribution by fluctuations of the concentration field itself, whereas \tilde{h}_r is caused by oscillations of the reaction sheet.

Variations of the \tilde{h} 's as functions of ω are plotted in Fig. 4, where solid and dotted curves, respectively, denote the near-equilibrium ($Da/Da_E = 5.12$), and the near-extinction ($Da/Da_E = 1.02$) cases. The upper figure plots

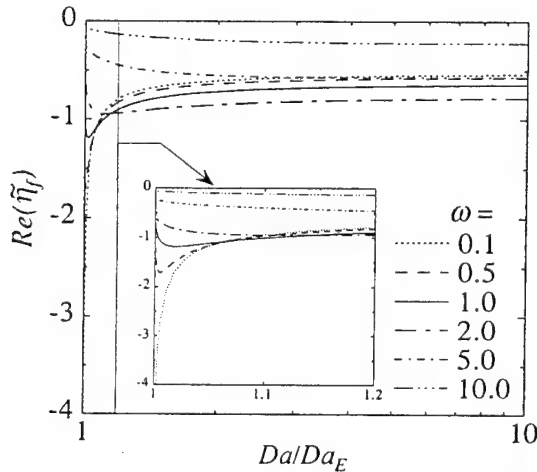


Fig. 3. Real parts of the magnitudes of reaction-sheet fluctuations as functions of steady Damköhler number for various frequencies.

the behavior in a phase plane of real and imaginary parts, showing the variation of the absolute magnitude as well as the phase angle of the \tilde{h} 's as a function of frequency. The arrows on the curves indicate the direction of the frequency increase. The lower plot of Fig. 4 is the projected view of the real part of the \tilde{h} 's with varying frequency along the y axis. To extract information from Fig. 4, first we select a value of ω on the y axis of the lower plot, then move horizontally to obtain the real value of the \tilde{h} 's. Subsequently by moving vertically to the corresponding curves in the upper plot, the magnitude and the phase can be obtained.

As observed from Fig. 3 that the reaction-sheet response is such that the reactivity becomes weaker as the strain rate increases with time during the oscillatory cycle, the upper plot of Fig. 4 shows that the \tilde{h}_r 's for both near-equilibrium and near-extinction cases take negative real values for low frequencies. On the other hand, the \tilde{h}_c 's for both cases have positive real values and are therefore in phase with the strain rate oscillation, in the sense of Rayleigh's criterion, because the net reactant flux at a fixed location of the mean reaction

sheet is directly related to the strain rate imposed at the boundary. As the frequency increases, each of the heat release responses spirals around the phase plane, showing the increase of phase lag and the variation of the magnitudes.

Next we observe the behavior of only the real parts of the \tilde{h} 's, shown in the lower plot of Fig. 4. In interpreting the results, we first recognize that ω is the ratio of the characteristic diffusion time to the characteristic oscillation time. That is, when $\omega \ll O(1)$, the burning rate is expected to respond quasi-steadily to the imposed oscillating strain rate because the characteristic oscillation time is much longer than that of diffusion in the flame structure. Consequently, the magnitudes of the \tilde{h} 's are all directly proportional to the amplitude of the strain rate oscillation, and are independent of the increase in ω . As ω approaches $O(1)$ value, the unsteady time becomes comparable to the diffusion time. Thus there is a lag between the imposed strain-rate variation and the response of the reaction sheet. For example, with an increase in the strain rate, the flame will not move to the new equilibrium position instantaneously. The reaction zone therefore experiences a higher concentration gradient which consequently leads to a higher burning rate. As ω continuously increases to very large values, however, the flame eventually stops responding to the oscillating strain rate as it approaches the steady-state limit.

We now consider the near-equilibrium flames in Fig. 4. Here starting with $\omega \ll 1$, increasing ω leads to an increase in \tilde{h}_c due to the increase in the instantaneous concentration gradient. The \tilde{h}_r also suffers a larger reduction in its response because the increased gradient causes more incomplete reaction. However, these amplifying behaviors are arrested and reversed around $\omega = O(1)$ with further increase in ω , producing the extrema in the respective responses. Near extinction, extrema are not observed because the reaction zone can respond more readily with increasing ω , thereby minimizing the aforementioned effects of amplification around $\omega = O(1)$. Consequently, the magnitudes of \tilde{h}_c and \tilde{h}_r decrease monotonically with increasing ω , approaching the quasi-steady limit for large ω .

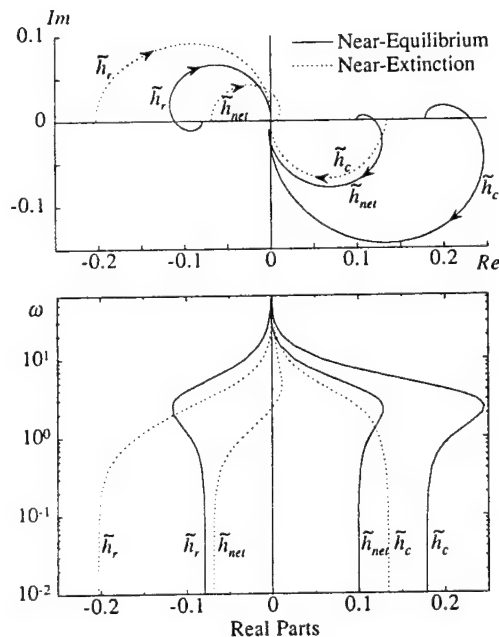


Fig. 4. Variations of heat release fluctuations as functions of frequency for near-equilibrium case and near-extinction case.

Based on the above understanding, the net response \tilde{h}_{net} can now be assessed. The lower plot of Fig. 4 shows that, in the near-equilibrium case, \tilde{h}_c governs the behavior of \tilde{h}_{net} , so that the net amount of heat release fluctuation \tilde{h}_{net} is in phase with the strain rate oscillation for all frequencies. In the near-extinction case, however, negative real values of \tilde{h}_r become dominant in the \tilde{h}_{net} behavior, reflecting the fact that response of the reaction rate to strain rate oscillation becomes more sensitive near extinction, as shown in Fig. 3. Consequently, for frequencies around $O(1)$ or less, the net burning rate response for the near-extinction flame could be out-of-phase with the imposed strain rate, implying that the net burning rate actually decreases as the blowing at the boundary increases during the unsteady oscillation. This contrasts with the common understanding of diffusion flame that the burning rate usually increases with the increase of reactant supply, based on the Burke-Schumann limit concept.

We next note that, from the standpoint of Rayleigh's criterion in acoustic instability, since acoustic velocity is $\pi/2$ phase shifted from acoustic pressure, the imaginary part of \tilde{h}_{net} is important to determine whether acoustic pressure will be amplified or attenuated by the acoustic velocity response. Figure 5 shows that $Im(\tilde{h}_{net})$ is negative for most of the range of Damköhler numbers, while it becomes large

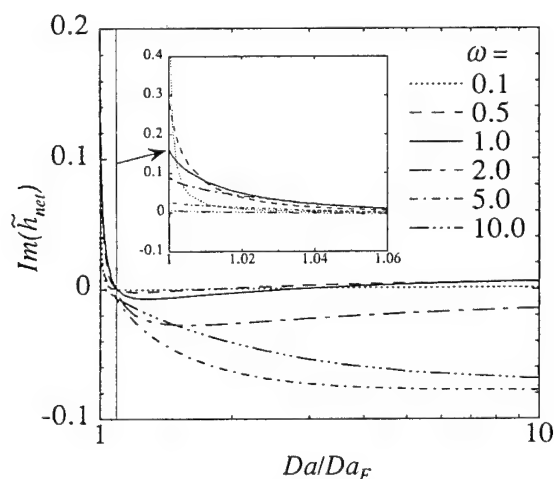


Fig. 5. Imaginary parts of the heat release fluctuations as functions of steady Damköhler number for various frequencies.

positive in the near-extinction case, suggesting the possibility that the existence of a sufficient number of near-extinction flamelets may change the acoustic instability of a rocket engine. Because an acoustic velocity cannot be uniquely determined unless the whole acoustic pressure field is known, inviscid considerations to relate the acoustic pressure with the unsteady strain rate should be pursued to better estimate the contribution of the velocity effect to acoustic amplification.

Finally, the effect of Lewis number of the fuel on the net heat release fluctuation is shown in Fig. 6. A previous steady-state analysis [11] has shown that the flame-sheet temperature can vary with strain rate depending on the Lewis number, which suggests that this influence will be carried over to the unsteady flame responses. Figure 6 demonstrates that this is indeed the case, with the response of \tilde{h}_{net} in general more significant for small Le_F , because of the relatively higher reactant diffusivity.

CONCLUSIONS

In the present paper, we have studied the response characteristics of a counterflow dif-

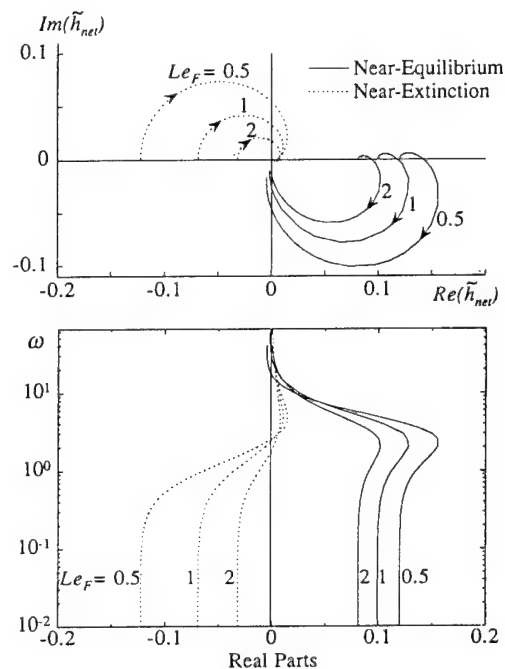


Fig. 6. Effect of the Lewis number of fuel on the net heat release fluctuations as functions of frequency.

fusion flame to a periodically oscillating strain rate, using large activation energy asymptotics. In particular, the responses of the reaction sheet as well as the heat release rate, in terms of their magnitude and phase, have been investigated for the near-extinction and the near-equilibrium flames. As suggested by the static S-curve behavior, the reaction-sheet response was found to be more sensitive to the flow-field unsteadiness when the steady flame is closer to the extinction condition. Consequently, the heat release behavior of the near-extinction flame is dominated by the reaction-sheet response rather than by the otherwise important fluid-dynamic effects, resulting in a significant difference in the heat release response of the two flames. It was also observed that there exist a nonmonotonic behavior in the heat release response of the near-equilibrium flame, consistent with a previous study [7], confirming that the flame behavior is mainly controlled by fluid dynamics when chemistry is sufficiently fast. Results of the present study suggest the possibility of a qualitative change in the characteristics, such as the stability response, of the near-extinction flame from that of the conventional flame in the Burke-Schumann limit. Lastly, the effect of Lewis number on the unsteady heat release response was shown to be consistent with our previous understanding.

As application to combustion instabilities in aero and rocket engines, the present linear perturbation analysis for the velocity response of the diffusion flame can be combined with the previous analysis on the pressure response [6] by superposition principle, which will then provide complete information of the flame response to the acoustic perturbation.

We finally comment that the present analysis is restricted to flames with sufficiently thin reaction zone with a single time scale. If detailed chemistry is considered, different flame

structures may emerge. For instance, the hydrogen/air diffusion flame near extinction consists of a radical recombination zone as thick as the diffusive transport layer so that the quasi-steady reaction zone is no longer valid. In such a situation, direct coupling of unsteadiness and recombination reaction may lead to an even larger finite-rate chemistry effect. More advanced analytical study for such flames can perhaps be achieved by the use of reduced mechanisms.

This research was supported by the U. S. Air Force Office of Scientific Research through grants awarded to Princeton University and the University of California at San Diego.

REFERENCES

1. Haworth, D. C., Drake, M. C., Pope, S. B., and Blint, R. J., *Twenty-Second Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1988, pp. 589-597.
2. Stahl, G., and Warnatz, J., *Combust. Flame* 85:285-299 (1991).
3. Darabiha, N., *Combust. Sci. Technol.* 86:163-181 (1992).
4. Ghoniem, A. F., Soteriou, M. C., and Knio, O. M., *Twenty-Fourth Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1992, pp. 223-230.
5. Liñán, A., and Williams, F. A., *Fundamental Aspects of Combustion*, Oxford University Press, 1993, Ch. 5.
6. Kim, J. S., and Williams, F. A., *Combust. Flame* (in press).
7. Strahle, W. C., *Tenth Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1965, pp. 1315-1325.
8. Williams, F. A., *AIAA J.* 3:2112-2124 (1965).
9. McIntosh, A. C., *Combust. Sci. Technol.* 75:287-309 (1991).
10. Liñán, A., *Acta Astronaut.* 1:1007-1039 (1974).
11. Law, C. K., and Chung, S. H., *Combust. Sci. Technol.* 29:129-145 (1982).

Received 1 December 1993; revised 20 April 1994

Comments

D. Lozinski, *University of Illinois, USA*. Could you please comment on the anticipated stability of your model's solution. Specifically, Peters [1] demonstrated that Linan's premixed regime is unstable to a fast time instability when the

Lewis number (Le) is 1. Stewart [2] determined that the fast time instability could arise when $Le > 1$ with no heat loss (your $m = 0$). However, our recent work on smouldering required a re-examination of the fast time instability [in

preparation] where we found $Le < 1$ has a stabilizing effect even when heat losses are present.

REFERENCES

1. N. Peters, *Comb. Flame*, 33:315 (1978).
2. D. S. Stewart, *Comb. Flame*, 64:157 (1986).

Authors' Reply. There is some controversy about the interpretation of the results of Peters [1] that you cite, in particular concerning the form of expansion required and the need to match to possible perturbations in outer zones when m is near zero. It is interesting that similar questions arise even for stability of diffusion flames in the diffusion-flame regime (besides the premixed-flame regime addressed in the paper and your comment) at these shorter time scales, as we are finding in work yet unpublished, where we too conclude that Lewis numbers less than unity have a stabilizing effect. However, as discussed in the written paper and more fully in its Ref. 6, these time scales are shorter than those commonly encountered in turbulent flames and acoustic instability, and therefore we have not analyzed that high-frequency regime in the present paper.

A. Ghoniem, MIT, USA. In the previous symposium, we presented a numerical study of a similar problem in which we showed that the flame is most responsive to oscillations when

their mean is close to the steady extinction strain, i.e., when chemical rates are important, with their concomitant non-linear dependence of the gas dynamic variables. In a linearized theory like yours, do you expect to recover this effect?

Authors' Reply. We are aware of your paper in the previous Symposium and in fact referenced that, in the written paper, as a source of references to previous works on the subject. Our linear theory of course does not include non-linear effects but surely does show that the flame is most responsive close to steady extinction, as fully demonstrated in the written paper.

G. Kosaly, University of Washington, USA. Your first transparency shows that you assume that the dynamics of the molecular mixing is quasi-steady. This assumption may break down at high frequencies. Have you looked into this possibility?

Authors' Reply. On the contrary, the analysis accounts for unsteady molecular mixing but assumes that the chemistry in the reaction zone is quasi-steady. This assumption, of course, breaks down at sufficiently high frequencies, but these frequencies are higher than those commonly encountered, and we have not analyzed that regime. More information on this can be found in the written paper and especially in Ref. 6.

Thermophoretic Effects on Seeding Particles in LDV Measurements of Flames

C. J. SUNG and C. K. LAW *Department of Mechanical and Aerospace Engineering, Princeton University, Princeton, NJ 08544*
and R. L. AXELBAUM *Department of Mechanical Engineering, Washington University, St. Louis, MO 63130*

(Received May 5, 1993)

ABSTRACT—The motion of LDV seeding particles under the influence of viscous and thermophoretic forces in the rapidly-accelerating, high-temperature-gradient flame environment was studied via the counterflow premixed twin-flame configuration. Results demonstrate that thermophoretic force can induce significant lag between the fluid and particle velocities in the active preheat zone of a flame, and suggest that caution should be exercised when interpreting LDV data obtained in this region. A thermophoretic velocity correction to the LDV-determined velocity, with known experimental or computational temperature profile, is proposed. Additional considerations of LDV diagnostics and its determination of laminar flame speed are also presented.

INTRODUCTION

The application of laser-Doppler velocimetry (LDV) for velocity measurements in flames and the imaging of reacting flows by laser sheet light scattering require the introduction of seeding particles as light scatterers. The accuracy of these measurements therefore depends critically on how closely the seeding particles follow the flow. In addition to viscous drag, which causes the particles to follow the fluid motion, there are other forces which can cause the motion of the seeding particles to depart from it. Examples are the electrostatic, gravitational, centrifugal, acoustic, diffusiophoretic, photophoretic and thermophoretic forces (cf. Durst *et al.*, 1981; Gomez and Rosner 1993). Among them, thermophoresis is of particular significance for flame-related measurements (Rosner *et al.*, 1992; Gomez and Rosner 1993) because of the presence of high temperature gradients in the narrow thermal expansion region. Since the particles experience a thermophoretic force in the direction opposite to that of the temperature gradient, the velocity measured by LDV can be higher or lower than the actual value depending on the direction of the flow relative to that of the temperature gradient (Talbot *et al.*, 1980). Similarly, for instantaneous flame surface imaging techniques, the information obtained from transient light scattering can be used, without correction, only if the so-called thermophoretic displacement criterion is met in addition to the more obvious criterion of characteristic particle stopping time (Gomez and Rosner 1993).

Recently this concern with LDV became apparent in studies involving counterflow diffusion and premixed flames (Chelliah *et al.*, 1991; Sung and Law 1993). Here the LDV measurements agreed well with the computed values in the cooler, decelerating part of the flow upstream of the flame, but were found to significantly lag the calculated values in the rapidly-accelerating preheat region of the flame in which substantial

thermal expansion occurs over a very short distance. Estimates (Sung and Law 1993) showed that the difference could be satisfactorily accounted for by the influence of the thermophoretic force on the seeding particles.

The objective of the present study is to further explore the importance of thermophoretic force on particles in the rapidly-accelerating, high-temperature-gradient environment of flames by analyzing in detail the dynamics of the seeding particles in the counterflow flames of Sung and Law (1993). The advantage of using these particular flames is that for the present purpose they have been completely characterized experimentally as well as computationally. Specifically, profiles of the velocity, temperature, and major species concentrations across the flame have been determined by using non-intrusive laser diagnostic techniques, while the flame structure has also been computationally simulated with detailed transport and chemistry. Furthermore, the measured and computed values agree closely, except for the velocity lag in the rapidly accelerating region just mentioned. This well-characterized flame environment therefore allows accurate calculations of the various transport coefficients needed for the determination of the particle dynamics.

In the following we shall first present formulation of the equation of particle motion, which is followed by the solution and discussion of the results. Additional considerations of LDV diagnostics will also be presented.

FORMULATION

The equation of motion for a spherical particle relative to an infinite, stagnant, viscous fluid was first derived by Basset (1888). For a moving fluid, Basset's equation can be expressed as (Hjelmfelt and Mockros 1966; Durst *et al.*, 1981):

$$\begin{aligned} \frac{\pi d_p^3}{6} \rho_p \frac{dv_p}{dt} = & -3\pi\mu d_p(v_p - v_F) + \frac{\pi d_p^3}{6} \rho_F \frac{dv_F}{dt} - \frac{1}{2} \frac{\pi d_p^3}{6} \rho_F \frac{d(v_p - v_F)}{dt} \\ & - \frac{3}{2} d_p^2 (\pi\mu\rho_F)^{1/2} \int_{t_0}^t \frac{d(v_p - v_F)}{d\xi} \frac{d\xi}{(t - \xi)^{1/2}} \end{aligned} \quad (1)$$

Accelerating force
Stokes drag force
Pressure gradient force on fluid
Fluid resistance to accelerating sphere
Drag force associated with unsteady motion

where d_p is the particle diameter, v the velocity, ρ the density, μ the fluid viscosity, t the time, and subscripts "P" and "F" respectively designate the particle and the fluid. External forces, such as gravitational, centrifugal and electrostatic, are neglected. Since the densities of the solid particles used for seeding flames are normally much greater than that of the gas mixture, terms involving ρ_F in Equation (1) can be neglected compared with those involving ρ_p . The particle motion therefore is governed by the first two terms in Equation (1), and corresponds to the Type III approximation of Hjelmfelt and Mockros (1966). Furthermore, since the no-slip boundary condition ceases to be accurate for the small, sub- μm particles used in seeding, a slip correction factor is needed to modify the Stokes' law. The Knudsen-Weber form of the slip

correction factor C for all Knudsen number, Kn , is given by

$$C = 1 + Kn[\alpha + \beta \exp(-\gamma/Kn)], \quad (2)$$

where α , β , and γ are characteristic parameters determined by fitting the Knudsen-Weber formula to the experimental data. For solid particles these values are 1.142, 0.558, and 0.999 respectively (Allen and Raabe 1985). Furthermore, the particle Knudsen number is defined as

$$Kn = \frac{2\lambda}{d_p}, \quad (3)$$

where λ , the mean free path of the gas molecules, is related to the coefficient of viscosity by (cf. Kennard 1938; Hirschfelder *et al.*, 1954)

$$\mu = \Phi \rho_F \lambda c. \quad (4)$$

Here c is the mean velocity of the gas molecules and Φ is a constant which approaches 0.491 for repulsive intermolecular forces. Thus the Stokes drag force on a particle with slip is given by

$$F_{SD} = \frac{-3\mu d_p (v_p - v_F)}{C}. \quad (5)$$

In the near-continuum regime, the thermophoretic force on a spherical particle due to temperature gradient ∇T was obtained by Brock (1962) (cf. Talbot *et al.*, 1980) as

$$F_{TP} = \frac{-6\pi\mu\eta d_p C_s \left(\frac{k_F}{k_p} + C_t Kn \right) \frac{\nabla T}{T}}{(1 + 3C_m Kn) \left(1 + 2\frac{k_F}{k_p} + 2C_t Kn \right)}, \quad (6)$$

where $\eta = \mu/\rho$, k is the thermal conductivity, and C_m , C_s , and C_t are respectively the momentum exchange, thermal slip, and thermal exchange coefficients specified by the kinetic theory of gases. Talbot *et al.* (1980) suggested $C_m = 1.14$, $C_s = 1.17$, and $C_t = 2.18$ such that this fitting formula appears to be reasonably satisfactory for the entire range of Knudsen numbers. We also note that particle nonsphericity could in general play an important role in determining the thermophoretic properties (cf. Williams 1986; Williams 1987; Garcia-Ybarra and Rosner 1989). For instance, an elongated particle will thermophoretically drift faster than a sphere when it is oriented parallel to $-\nabla T$ (Garcia-Ybarra and Rosner 1989). The present assumption of particle sphericity is justified by the close agreements between the predicted and experimental results to be presented later.

Combining the above, the particle motion is governed by

$$m_p \frac{dv_p}{dt} = F_{SD} + F_{TP}, \quad (7)$$

where $m_p = \rho_p \pi d_p^3/6$ is the particle mass. This is the equation of motion for the seeding particles.

In Sung and Law (1993), numerical calculations of the symmetric counterflow, premixed, nitrogen-diluted methane/air flames were performed with potential flow boundary conditions and detailed reaction mechanisms and transport properties. The reaction mechanism used was a complete C_2 scheme consisting of 28 species and 151 elementary reaction steps. The subroutine library CHEMKIN was used to evaluate the thermodynamic properties of the mixture (Kee *et al.*, 1980). In conjunction with CHEMKIN, a transport package provided the transport properties of the mixture (Kee *et al.*, 1983), with the combination averaging formula and the Wilke formula used to respectively evaluate the thermal conductivity and viscosity of the mixture. The alumina seeding particles used in the experiments had a density of 3.97 g/cm^3 and a nominal diameter of $0.3 \mu\text{m}$. The variation of particle conductivity with temperature was fitted from standard tables (Rohsenow *et al.*, 1985). Using local values of the mixture and the particle, and appropriate interpolation between values at mesh points, the finite-difference form of Equation (7) was solved for v_p .

RESULTS AND DISCUSSIONS

The gaseous environment in which the particles traverse is a nitrogen-diluted methane/air flame ($\phi = 0.95$, $N_2/O_2 = 5$) at one atmosphere pressure and 300 K upstream temperature. We first consider only the viscous drag force F_{SD} in Equation (7). Figure 1 shows the calculated particle motion for different particle sizes with the initial condition $v_p = v_F$. The strain rate κ , defined as the negative of the axial velocity gradient just upstream of the thermal mixing layer, is 568 sec^{-1} for this strongly-strained situation. The local Reynolds number based on the relative velocity between the particle and the fluid, $Re = \rho_F d_p (v_p - v_F) / \mu$, was also calculated to ensure

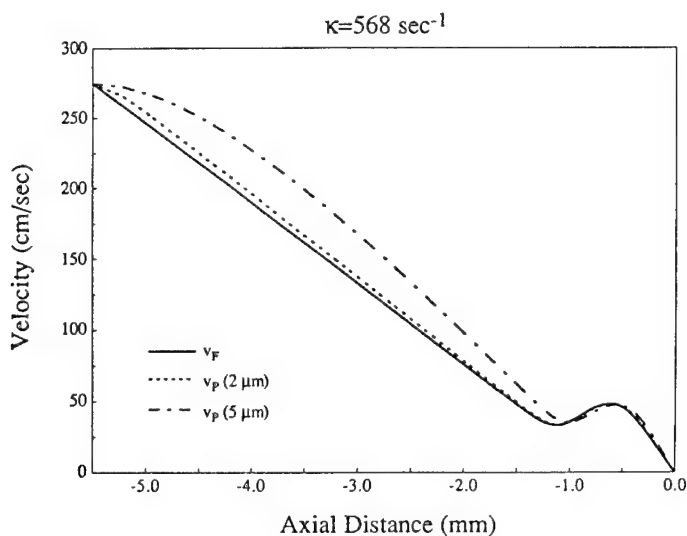


FIGURE 1 Solutions of particle motion under the influence of the viscous drag force for various particle sizes in a given counterflow flame ($\kappa = 568 \text{ sec}^{-1}$).

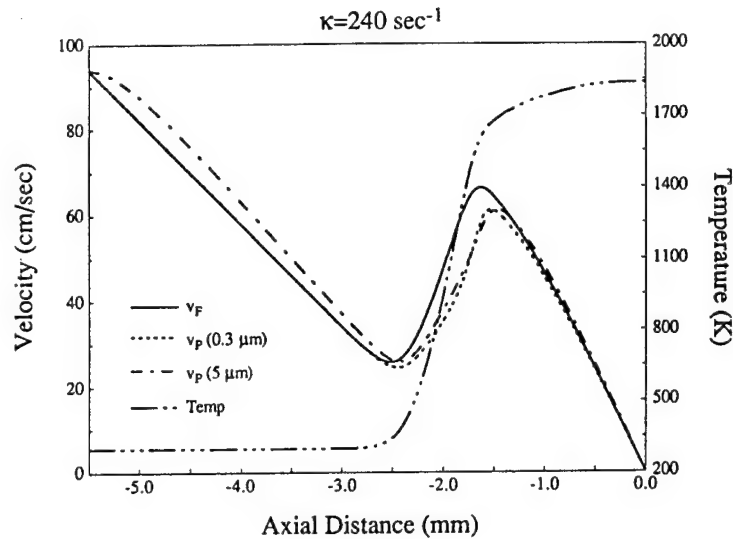


FIGURE 2 Solutions of particle motion under the influence of the viscous drag F_{SD} and the thermophoretic forces F_{TP} for various particle sizes in a given counterflow flame ($\kappa = 240 \text{ sec}^{-1}$).

that its value is less than unity such that Stokes drag law is applicable. It is seen that, for this strain rate, seeding particles with diameters greater than $2 \mu\text{m}$ substantially lag the gas flow. However, particles smaller than, say, $1 \mu\text{m}$ have been found to readily follow the flow both ahead of and inside the flame.

Figure 2 shows the particle motion in the presence of both drag and thermophoretic forces, with $\kappa = 240 \text{ sec}^{-1}$, for which experimental data exist. For the $0.3 \mu\text{m}$ particle, which readily follows the flow when subjected only to the drag force, significant velocity lag develops in the preheat zone when thermophoresis is considered. In the worst situation the lag is more than 15 cm/sec , which corresponds to a local v_F of about 60 cm/sec . This clearly demonstrates that thermophoretic effects should be considered in LDV measurements of flame dynamics and structure, especially when the measurement is conducted in the rapidly-accelerating thermal expansion region. It is also of interest to note that although particles with $d_p > 2 \mu\text{m}$ fail to follow the flow well ahead of the thermal mixing layer because of their inertia, the particle motion in the flame zone is however similar for particle sizes even up to $5 \mu\text{m}$. This is due to the combined effects of drag and thermophoretic forces, as will be discussed later.

In Figures 3 and 4 we compare the calculations with the available LDV data of sung and Law (1993) for $\kappa = 240$ and 348 sec^{-1} respectively. In performing these calculations we have noted that while agglomeration could have caused the actual particles in the flow to be larger than the individual particles, their exact size is not essential as long as d_p is less than $2 \mu\text{m}$, as shown in Figures 1 and 2. Thus we have used $d_p = 0.3 \mu\text{m}$ in the calculations. It is seen that the calculated values agree well with the experimental data, hence again demonstrating the importance of thermophoresis.

It is of interest to correct the measured LDV data so that the flow velocity can be accurately evaluated. Since particles of sufficiently small sizes, and hence small inertia,

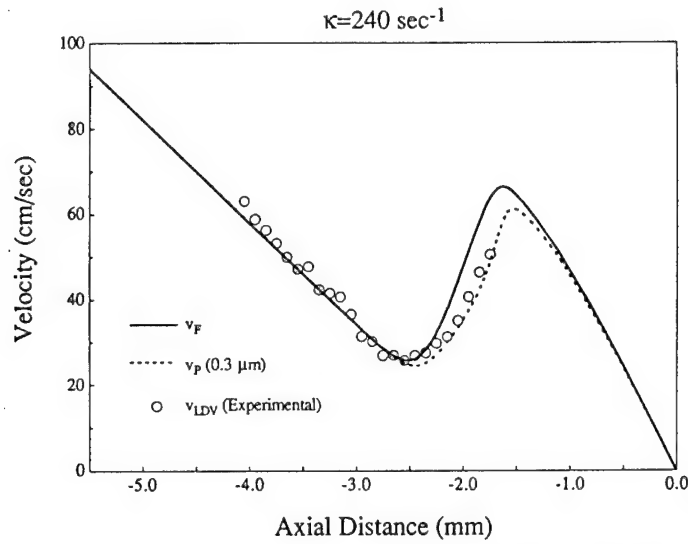


FIGURE 3 Comparison between the experimental and computed velocity profiles, for $\kappa = 240 \text{ sec}^{-1}$.

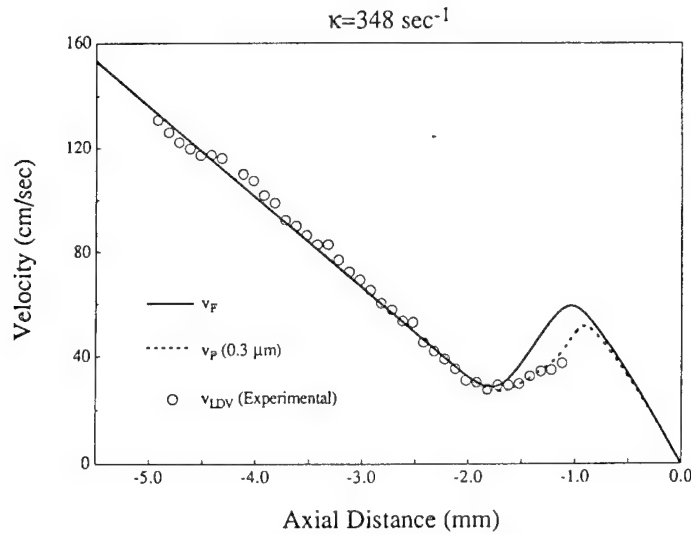


FIGURE 4 Comparison between the experimental and computed velocity profiles, for $\kappa = 348 \text{ sec}^{-1}$.

can readily adjust their motion in accordance with the local force field, it is reasonable to expect that the net force acting on them are much smaller than the individual forces due to either drag or thermophoresis, that is $|m_P(dv_P/dt)| \ll |F_{SD}|$ and $|m_P(dv_P/dt)| \ll |F_{TP}|$. From Equation (7) we can therefore make a "steady-state" approximation by setting $F_{SD} = -F_{TP}$. Using Equations (5) and (6) for these two forces then readily yields

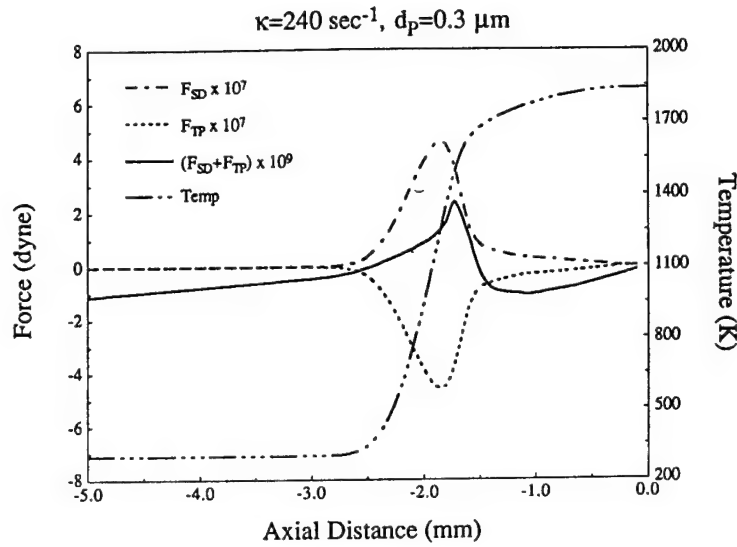


FIGURE 5 Comparison between F_{SD} , F_{TP} , and $F_{SD} + F_{TP}$, for $d_p = 0.3 \mu\text{m}$ and $\kappa = 240 \text{ sec}^{-1}$.

an expression for the actual flow velocity as

$$v_F = v_p + v_{TP} \quad (8)$$

where the thermophoretic velocity v_{TP} is given by

$$v_{TP} = \frac{-C}{3\pi\mu d_p} F_{TP} \quad (9)$$

Equations (8) and (9) allow approximate extraction of the flow velocity v_F from the LDV measurements of the particle velocity v_p . It may be noted that this steady-state assumption was implicitly used by Brock (1962) and Talbot *et al.* (1980), who provided the same expressions for the thermophoretic velocity.

The above steady-state approximation is expected to be adequate for particle sizes which are smaller than the maximum acceptable particle diameter for given velocity and temperature gradients. This is demonstrated for $d_p = 0.3$ and $1 \mu\text{m}$ in Figures 5 and 6 respectively, which show that $(F_{SD} + F_{TP})$ is about two orders smaller than either F_{SD} or F_{TP} . Figure 7, for $d_p = 5 \mu\text{m}$, shows that the accuracy of this approximation decreases for larger particles because they fail to closely follow the flow even without considering F_{TP} .

An alternate approach to determine the thermophoretic velocity is to experimentally measure the thermophoretic diffusivity, D_{TP} , in high temperature combustion environments, as was done by Gomez and Rosner (1993) for the D_{TP} of titanium dioxide particle by applying the phase separation phenomenon. The local particle thermophoretic velocity v_{TP} is then simply equal to $D_{TP} \nabla T/T$, and the local flow velocity can

be again corrected by using Equation (8). Since titanium dioxide particle is quite often used in seeding flames and its experimentally-determined D_{TP} is available, it is of interest to roughly estimate the maximum v_{TP} if the alumina particles of $0.3\text{ }\mu\text{m}$ nominal diameter in the experiments of Sung and Law (1993) were replaced by titanium dioxide particles. For the case of $\kappa = 240\text{ sec}^{-1}$ with $(\nabla T)_{\max} \approx 2000\text{ K/mm}$ at $T \approx 1300\text{ K}$, applying the result of Gomez and Rosner (1993), $D_{TP} \approx 0.5\eta$, which is applicable to particle sizes covering a diameter range between 2 nm and $0.4\text{ }\mu\text{m}$, and assuming $\eta_{\text{mixture}} \approx \eta_{\text{air}}$, the maximum v_{TP} is estimated to be about 14 cm/sec . This further demonstrates the importance of thermophoresis in combustion-related LDV measurements.

The above results demonstrate that if both temperature and velocity can be experimentally measured, the actual flow velocity profile can be evaluated by correcting the LDV-determined velocity by the corresponding local thermophoretic velocity v_{TP} . This approach is of course also applicable if computed rather than measured temperature profile is instead available. Since the need for the temperature profile is only for a reasonable evaluation of the transport properties and the temperature gradients, only an approximate profile would suffice for this purpose. For example, we have found that the temperature profile determined for the adiabatic one-dimensional flame is quite adequate for the evaluation, by overlaying the point of initial temperature rise of the one-dimensional flame at the minimum velocity point of the counterflow flame.

It is also of interest to note that, while the two competing forces, F_{SD} and F_{TP} , as well as the net force ($F_{SD} + F_{TP}$) increase with increasing particle size within the thermal mixing layer, as shown in Figures 5–7, the acceleration of the particle, dv_p/dt , have

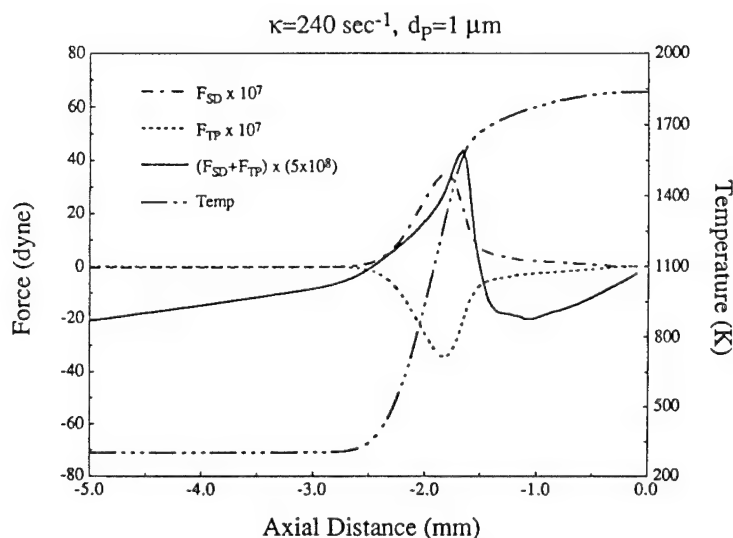


FIGURE 6 Comparison between F_{SD} , F_{TP} , and $F_{SD} + F_{TP}$, for $d_p = 1\text{ }\mu\text{m}$ and $\kappa = 240\text{ sec}^{-1}$.

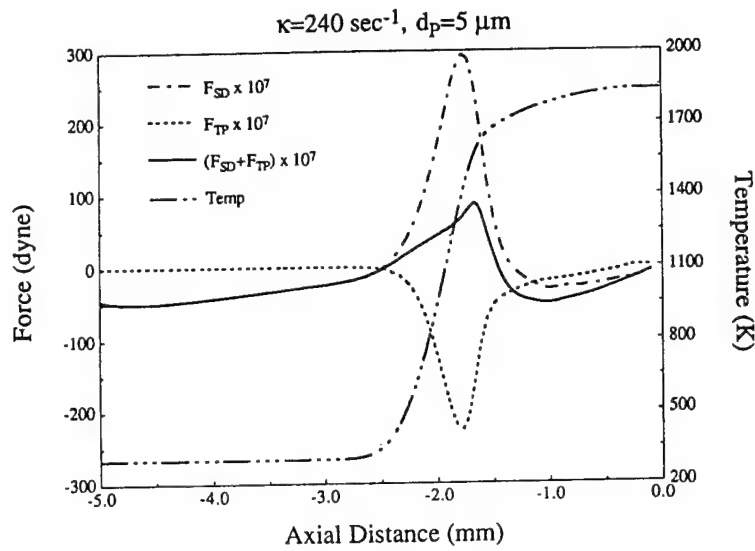


FIGURE 7 Comparison between F_{SD} , F_{TP} , and $F_{SD} + F_{TP}$, for $d_p = 5 \mu\text{m}$ and $\kappa = 240 \text{ sec}^{-1}$.

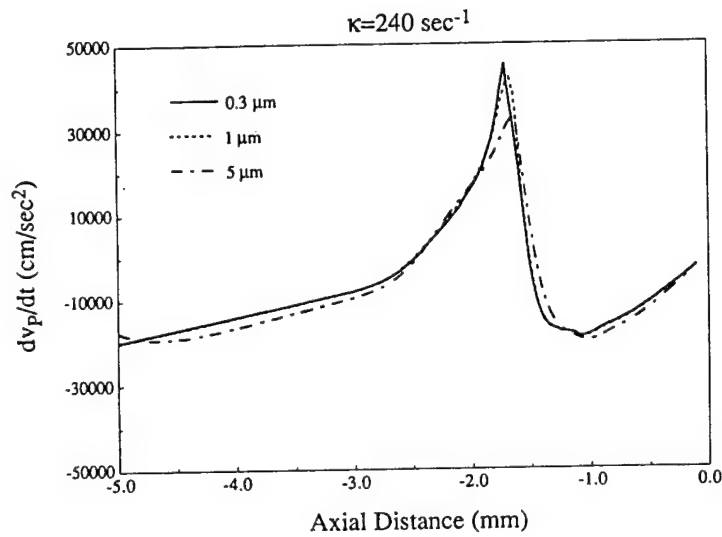


FIGURE 8 Comparison of the particle acceleration, dv_p/dt , for various particle sizes in a given counterflow flame ($\kappa = 240 \text{ sec}^{-1}$).

similar magnitudes for the diameter range of particle sizes studied herein. This point is demonstrated in Figure 8, and would explain the result of Figure 2 that particles of 0.3 and 5 μm closely follow each other in the thermal expansion zone.

In closing this section, it is therefore suggested that, for flame-related LDV measurement, we should first choose the seeding particle which is small enough to have a good

responsivity to a given velocity gradient in the uniform temperature situation. Subsequently, when interpreting the experimental data within the rapidly-accelerating thermal mixing layer, the thermophoretic correction proposed by Equation (8) can be applied to substantially improve the accuracy of the LDV-determined data.

ADDITIONAL CONSIDERATIONS

There are several ancillary points worth further discussion. First, we consider the maximum particle size for which the steady-state approximation is applicable to turbulent flows. Here we note that for LDV applications in uniform temperature turbulent flows, it is convenient to specify the desired amplitude response for a given turbulence frequency, from which the maximum acceptable particle diameter can be deduced. Hjelmfelt and Mockros (1966) solved the equation of particle motion, Equation (1), by expressing v_p and v_F as Fourier integrals. Their solution shows that, in order to have a frequency response of 1 kHz, the maximum acceptable particle diameters for TiO_2 in air and MgO in a post methane/air flame, with an assumed uniform temperature of 1800 K, are 1.3 and 2.6 μm respectively (cf. Durst *et al.*, 1981). It was further concluded that a diameter of 1 μm is an appropriate particle size to be used in a wide variety of gas flows with turbulence frequencies exceeding 1 kHz.

In order to correlate the strain rate with the "turbulence frequency", we simply consider a stagnation flow field whose axial velocity is given by $v = v_s - \kappa x$, for which the integration constant v_s can be written as $v_s = \kappa x_s$ with x_s being the location of the stagnation surface where $v = 0$. This relation can be readily integrated to yield $v(t) = v_s \exp(-\kappa t)$. Further taking the Fourier transform of $v(t)$, which is assumed to be zero for $t < 0$, yields the frequency spectrum of the velocity variation with time, $V(\omega)$. This spectrum has a magnitude of $|V(\omega)| = v_s/(\kappa^2 + \omega^2)^{1/2}$ and a phase angle $\tan^{-1}(-\omega/\kappa)$. It indicates the frequency content of $v(t)$ and can be thought of as the amount of v per unit frequency increment. This also allows us to interpret $V(\omega)$ as an amplitude density (cf. Doebelin 1980). For situations where the physical domain has a fixed dimension, namely $x_s = \text{constant}$, then $|V(\omega)|$ is proportional to $\kappa/(\kappa^2 + \omega^2)^{1/2}$. If we define the characteristic turbulence frequency ω^* as the high-frequency cutoff occurring at half of the maximum spectrum amplitude, then $\omega^* = \sqrt{3}\kappa$. Thus, the corresponding ω^* for the case of $\kappa = 568 \text{ sec}^{-1}$ shown in Figure 1 is about 1 kHz. The result of Figure 1 also agrees with the solution of Hjelmfelt and Mockros in that, for good frequency response above 1 kHz, seeding particles should have diameters less than 1 μm . Thus, to correct for thermophoresis with Equations (8) and (9), particles of 1 μm or less should be used.

The responsivity of the particle motion in the stagnation flow can also be assessed by considering Equation (1) for the uniform temperature situation. Here

$$m_p \frac{dv_p}{dt} = \frac{v_F - v_p}{B}, \quad (10)$$

where $B = (3\pi\mu d_p)^{-1}$ is the particle mobility, with $C = 1$ for simplicity. Since our attention is restricted to the situation where $v_F \approx v_p \gg (v_F - v_p)$, then it is reasonable to

assume that

$$\frac{dv_P}{dt} \approx \frac{dv_F}{dt} = v_F \frac{dv_F}{dx} = v_F \kappa. \quad (11)$$

Thus

$$m_p B \kappa = \frac{(v_F - v_P)}{v_F}, \quad (12)$$

which is the fractional lag of the particle. Equation (12) yields the convenient result that the particle lag is governed by the dimensionless group $m_p B \kappa$. For example, a $1 \mu\text{m}$ seeding particle subjected to a strain rate of 800 sec^{-1} in air would lag the flow by about 1%, and would thus be an acceptable seeding particle for most counterflow flame studies.

The second point of interest is the potential error associated with velocity bias as a consequence of the nonuniform velocity profile of the counterflow flame within the finite probe volume, causing velocities to be weighted nonuniformly when measuring the average velocity. Specifically, if the particles were distributed uniformly, then the presence of a velocity gradient through the probe volume would cause the higher velocities to be weighted more than the lower velocities because of the larger flux associated with high velocity particles. Consequently, the measured average velocity will be greater than the velocity in the center of the probe volume. When there is a temperature gradient across the probe volume, as there is in the preheat zone, the flux of particles can also be influenced by the gradient in the particle number density. The significance of these issues will be briefly discussed as they pertain to the counterflow premixed flame.

For purposes of discussion it is useful to divide the flame into three distinct regions: Region 1 is the cold unburned side of the flame where the temperature is constant at the freestream value; Region 2 is the preheat zone where the temperature increases rapidly; and Region 3 is the burned side where the temperature is constant at the flame temperature. We will consider each region separately.

The probability that a Doppler signal will be detected at a location x is most simply expressed by

$$P(x) = \frac{N(x)v(x)I(x)}{\int N(x)v(x)I(x)dx}, \quad (13)$$

where N is the particle number density and I is the laser intensity distribution within the probe volume. For simplicity, we shall assume that I is constant, which will lead to an overestimate of the velocity bias. Assuming that data is averaged over sufficiently large times, we can integrate $P(x)v(x)$ over the probe volume to obtain the LDV measured velocity v_{LDV} by substituting in $v(x) = \kappa x$. The unbiased velocity is that at the center of the probe volume v_m . The results in Region 1 show that the greatest velocity bias occurs at the lowest velocity and that in the limit of $v \rightarrow 0$ we obtain $(v_{LDV} - v_m) = \kappa \delta / 6$ where δ is the probe diameter, $\lim_{v \rightarrow 0} v_{LDV} = 2\kappa \delta / 3$, and $\lim_{v \rightarrow 0} v_m = \kappa \delta / 2$. For $\delta = 0.03 \text{ cm}$ and $\kappa = 600 \text{ sec}^{-1}$, $\lim_{v \rightarrow 0} (v_{LDV} - v_m) = 3 \text{ cm/s}$. The

actual velocity bias will be considerably less than this because $v > 0$ and $I(x)$ is a Gaussian function, not a constant. Under most circumstances, then, velocity bias can be neglected in this region.

In Region 2, where the temperature and velocity are increasing, the velocity bias is even less than in Region 1 because the increase in v with x is offset by a nearly corresponding decrease in ρv (i.e., Nv). Estimates of the velocity bias in this region show that for typical flame studies, the bias is less than 1 cm/s which is within the experimental uncertainty of the LDV measurements.

Measurements in Region 3 have the greatest potential for error from velocity bias because the gradient in v is greatest and N is constant. However, dv/dx in this region is typically only 2–3 times that in Region 1, i.e., $\kappa_3 \approx 2\kappa_1$, so the bias can again be neglected except for regions close to the stagnation point where $v \rightarrow 0$ such that the error could be substantial and corrections due to velocity bias would be appropriate.

The third ancillary point concerns the various methodologies used to determine the laminar flame speeds of combustible mixtures. In particular, there has been discussions on the relative merits of determining these values based on the flow velocity either upstream or downstream of the preheat zone. For example, in the twin-flame counter-flow methodology, the former would correspond to identifying the reference velocity as the minimum in the velocity profile (Wu and Law 1985) while the latter the maximum (Mendes-Lopes *et al.*, 1983; Mendes-Lopes *et al.*, 1985). In light of the present understanding, it is clear that the thermophoretic force could significantly reduce the downstream reference velocity, thereby lead to lower values of the laminar flame speed eventually determined if thermophoretic correction is not made. The same concern also applies to using the flow velocity upstream of the luminous zone as a reference because significant thermal expansion has already taken place as the flow reaches the state of active chemiluminescence. It is therefore suggested that use of LDV instrumentation in flame measurements should be limited to the flow region upstream of the active preheat zone, unless thermophoretic effects are properly taken into account.

CONCLUDING REMARKS

The present study has yielded the following points of interest on the accuracy of LDV measurements, especially those related to stagnation flows and to flames.

1. For a uniform-temperature stagnation point flow, the strain rate κ can be defined as the characteristic turbulence frequency of the system. The responsiveness of the particle motion and the maximum acceptable particle diameter for a given turbulence frequency can be assessed by using either the solution of Hjelmfelt and Mockros (1966) or Equation (12) demonstrated herein. The results show that 1 μm would be an acceptable seeding particle size for most stagnation point flow studies of hydrocarbon/air flames. The particle size would need to be smaller for flames with much higher flame speeds such as those of hydrogen or enriched with oxygen.

2. Concerns with the potential error of velocity bias due to the finite probe volume as shown to be negligible for counterflow flames except for regions close to the stagnation point where $v \rightarrow 0$.
3. For flame-related LDV measurements, significant velocity lag develops within the rapidly accelerating thermal expansion region due to the presence of the high temperature gradient. The agreement between the available experimental data of Sung and Law (1993) on counterflow premixed flames and the present calculations demonstrate the importance of thermophoretic forces imposed on the LDV seeding particles.
4. If both temperature and velocity can be measured in an experiment, the actual flow velocity profile can be obtained by correcting the LDV-determined velocity with the corresponding local thermophoretic velocity v_{TP} given by Equation (9) or $D_{TP} \nabla T/T$ if the information on D_{TP} is available. This approach can improve the accuracy of the velocity measurement as long as the particle size is smaller than the maximum acceptable particle diameter for a given velocity gradient or characteristic turbulence frequency. If an experimental temperature profile is not available, a representative calculated profile can also be used.
5. When using the twin-flame counterflow methodology to determine the laminar flame speed of combustible mixtures, LDV measurements should be limited to the flow region upstream of the active preheat zone if thermophoretic correction is not made.

ACKNOWLEDGEMENTS

This work has been supported by the Air Force Office of Scientific Research and the Division of Basic Energy Sciences of the Department of Energy, under the technical monitoring of Dr. J. M. Tishkoff and Dr. O. P. Manley respectively. We appreciate the very helpful discussions with Professor P. Cho of Michigan Technological University and Dr. P. G. Felton of Princeton University. The contribution of Mr. J. A. Eng of Princeton University in the computations is also gratefully acknowledged.

REFERENCES

- Allen, M. D. and Raabe, O. G. (1985). *Aerosol Sci. Technol.* **4**, 269.
- Brock, J. R. (1962). *J. Colloid Sci.* **17**, 768.
- Chelliah, H. K., Law, C. K., Ueda, T., Smooke, M. D. and Williams, F. A. (1991). *Twenty-Third Symposium (International) on Combustion*, p. 503, The Combustion Institute.
- Doebelin, E. O. (1980). *System Modeling and Response: Theoretical and Experimental Approaches*, John Wiley & Sons.
- Durst, F., Melling, A., and Whitelaw, J. H. (1981). *Principles and Practice of Laser-Doppler Anemometry*, Academic.
- Garcia-Ybarra, P. and Rosner, D. E. (1989). *AIChE J.* **35**(1), 139.
- Gomez, A. and Rosner, D. E. (1993). *Combust. Sci. Technol.* **89**, 335.
- Hirschfelder, J. O., Curtiss, C. F. and Bird, R. B. (1954). *Molecular Theory of Gases and Liquids*, Wiley.
- Hjelmfelt, A. T., Jr. and Mockros, L. F. (1966). *Appl. Sci. Res.* **16**, 149.
- Kee, R. J., Miller, J. A. and Jefferson, T. H. (1980). *Sandia Report* 80-8003.
- Kee, R. J., Warnatz, J. and Miller, J. A. (1983). *Sandia Report* 83-8209.
- Kennard, E. H. (1938). *Kinetic Theory of Gases*, McGraw-Hill.
- Mendes-Lopes, J. M. C., Daneshyar, H. and Ludford, G. S. S. (1983). *Nineteenth Symposium (International) on Combustion*, p. 413, The Combustion Institute.
- Mendes-Lopes, J. M. C. and Daneshyar, H. (1985). *Combust. Flame*, **60**, 29.

- Rohsenow, W. M., Hartnett, J. P., and Ganic, E. N. (1985). *Handbook of Heat Transfer Fundamentals*, McGraw-Hill.
- Rosner, D. E., Mackowski, D. W., Tassopoulos, M., Castillo, J. L. and Garcia-Ybarra, P. (1992). *I&EC-Research (ACS)* **31**, 760.
- Sung, C. J. and Law, C. K. (1993). *AIAA Thirty-first Aerospace Sciences Meeting*, Paper No. 93-0246; also submitted to *Combust. Flame*.
- Talbot, L., Cheng, R. K., Schefer, R. W. and Willis, D. R. (1980). *J. Fluid Mech.* **101**, 737.
- Williams, M. M. R. (1986). *J. Physics D, Applied Physics*, **19**, 1631.
- Williams, M. M. R. (1987). *J. Physics D, Applied Physics*, **20**, 354.
- Wu, C. K. and Law, C. K. (1985). *Twentieth Symposium (International) on Combustion*, p. 1941, The Combustion Institute.

FURTHER CONSIDERATIONS ON THE DETERMINATION OF LAMINAR FLAME SPEEDS WITH THE COUNTERFLOW TWIN-FLAME TECHNIQUE

C. M. VAGELOPOULOS AND F. N. EGOLFOPOULOS

*Department of Mechanical Engineering
University of Southern California
Los Angeles, CA 90089-1453, USA*

AND

C. K. LAW

*Department of Mechanical and Aerospace Engineering
Princeton University
Princeton, NJ 08544, USA*

The accuracy of the laminar flame speed determination by using the counterflow twin-flame technique has been computationally and experimentally examined in light of the recent understanding that linear extrapolation of the reference upstream velocity to zero strain rate would yield a value higher than that of the laminar flame speed, and that such an overestimate can be reduced by using either lower strain rates and/or larger nozzle separation distances. A systematic evaluation of the above concepts has been conducted and verified for the ultralean hydrogen/air flames, which have relatively large Karlovitz numbers, even for small strain rates, because of their very small laminar flame speeds. Consequently, the significantly higher values of the previous experimentally measured flame speeds, as compared with the independently calculated laminar flame speeds, can now be attributed to the use of nozzle separation distances that were not sufficiently large and/or strain rates that were not sufficiently small. Thus, by using lower strain rates and larger nozzle separation distances, the experimentally and computationally redetermined values of these ultralean hydrogen/air flames agree well with the calculated laminar flame speeds. The laminar flame speeds of methane/air and propane/air mixtures have also been experimentally redetermined over extensive ranges of the equivalence ratio and are found to be slightly lower than the previously reported experimental values.

Introduction

Because of the fundamental and practical importance of the laminar flame speed, S_u^0 , commonly known as the laminar burning velocity, extensive efforts have been expended over the years toward achieving accurate determinations of their values. The early efforts, however, have resulted in data with substantial scatter, even for experiments apparently conducted with great care. Furthermore, the extent of scatter also did not improve with time, as demonstrated by the summary plot of Andrews and Bradley [1]. While it was recognized quite early that the laminar flame speeds could be substantially modified by aerodynamic stretch, with or without the additional influence of preferential diffusion [2], it was not until recently that systematic efforts have been mounted toward eliminating these stretch effects either during experimentation or through further data processing. A strong motivation for such efforts was the recent interest in *partially* validating chemical kinetic schemes through numerical simulation of the flame structure and hence the associated S_u^0 .

One of such efforts was the development of the counterflow twin-flame technique for the determination of S_u^0 [3,4]. Specifically, a twin-flame configuration is established in counterflow created by impinging two identical, nozzle-generated flows of the combustible mixture of interest onto each other. By mapping the flow field using laser Doppler velocimetry (LDV) and identifying the minimum in the velocity profile as a reference upstream burning velocity, $S_{u,ref}$, and the velocity gradient ahead of the minimum point as the strain rate experienced by the flame, K , $S_{u,ref}$ can be plotted for various K (see inset of Fig. 1 for definition). Based on the behavior of the experimental data and the results from theoretical studies, it was suggested that $S_{u,ref}$ should vary linearly with K , for sufficiently small values of K , such that a linear extrapolation to $K = 0$ should yield the S_u^0 of the unstrained flame (see Fig. 2 for representative extrapolations). The extrapolated experimental values thus determined appear to agree quite well with those obtained through independent numerical computation of the unstrained planar flame with detailed transport and chemistry, for systems whose

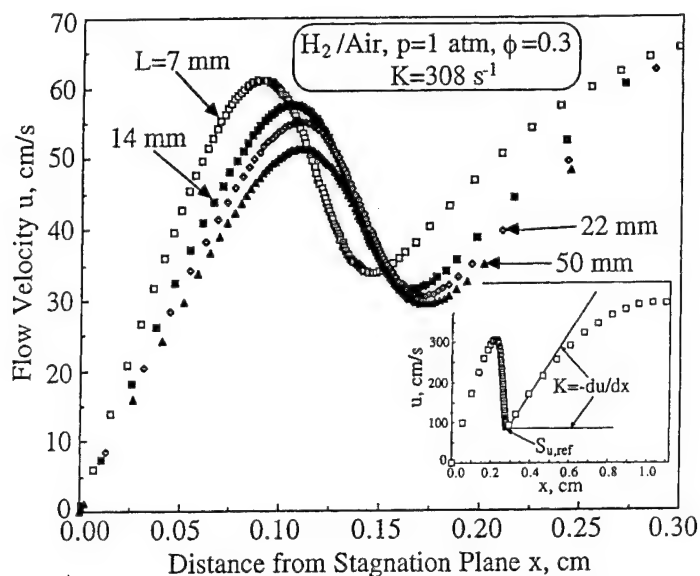


FIG. 1. Spatial variation of the numerically determined flow velocity profiles in the counterflow, for $\phi = 0.3$ atmospheric hydrogen/air mixture with $K = 308 \text{ s}^{-1}$ and $L = 7, 14, 22$, and 50 mm , and the definition of K and $S_{u,ref}$.

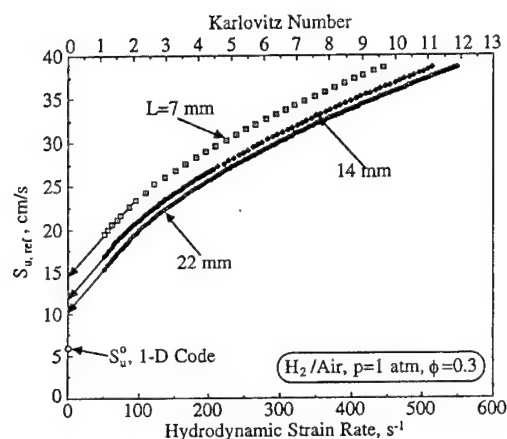


FIG. 2. Variation of the numerically determined $S_{u,ref}$ with K and Karlovitz number for $\phi = 0.3$ atmospheric hydrogen/air mixture and $L = 7, 14$, and 22 mm .

chemistry is considered reasonably well known as far as the computation of S_u° is concerned.

The technique discussed above is based on the assumption of a linear relation between $S_{u,ref}$ and K for small K . Tien and Matalon [5], however, subsequently showed through asymptotic analysis that the relation is actually not linear as $K \rightarrow 0$, and that a linear extrapolation would yield a slightly higher value than the true S_u° . This point was further substantiated by Dixon-Lewis [6] through detailed numerical simulation of the counterflow flame with a finite separation distance L between the nozzle exits. The difference observed [6], however, was again small in that, for stoichiometric methane/air flames,

the extrapolation yields $S_u^\circ = 39.6 \text{ cm/s}$, while direct computation yields 36.7 cm/s . The results of Ref. 6 also did not show any dependence on the separation distance of the nozzle.

Chao et al. [7] subsequently performed an asymptotic analysis of the counterflow problem, allowing for the plug flow boundary condition and a finite nozzle separation distance, L . The solution again yields the nonlinear variation as $K \rightarrow 0$, although it also showed that, for a given flame thickness, the overestimate can be reduced by either decreasing K or increasing L such that perturbation of the flow field by the flame is minimized. Note that, since a larger L implies the use of a proportionally larger nozzle if the ratio of L to the nozzle diameter is to be kept fixed, this requirement could have practical experimental limitations due to either inadequate flow rate or safety considerations when working with a large flow rate of combustible gas. Chao et al. [7] further performed detailed numerical simulation and experimentation with methane/air flames of selected stoichiometries. Results demonstrate that, contrary to the finding of Ref. 6, the overestimate indeed can be progressively reduced by using larger values of L . Furthermore, for the mixtures and hence the range of burning velocities studied, results obtained with nozzle separation distances of 14 and 22 mm show little difference, indicating that the use of L within this range should yield laminar flame speeds of reasonable accuracy, at least for methane/air mixtures.

The objectives of the present study are to further explore, computationally and experimentally, the influence of the separation distance on the accuracy of the laminar flame speed determination, and to re-determine the laminar flame speed values for the im-

portant fuel/air mixtures of hydrogen/air, methane/air, and propane/air. Based on the results of Chao et al. [7], we expect the revision from previously reported S_u^0 to be small for methane/air and propane/air flames. The revision, however, is needed because recent advances in the kinetic database require an increased accuracy for resolution. For hydrogen/air flames, we shall limit our study to ultra-lean mixtures for safety considerations. For these very weak flames, however, the influence due to the finite separation distance is also expected to be the greatest because they are most susceptible to stretch effects. We are especially interested in resolving the discrepancy observed in our previous work [8] that the experimentally measured S_u^0 for these flames were substantially higher than the calculated values based on apparently reliable hydrogen oxidation kinetics. In light of the new understanding that linear extrapolation tends to result in overestimates, it is possible that these previous experimental values could indeed be too high. Thus, for these flames, the revision is crucial because of the substantial discrepancy in the earlier data and because of its fundamental implications on the hydrogen oxidation kinetics.

The numerical and experimental methodologies will be presented in the next two sections, which will be followed by results on hydrogen/air, methane/air, and propane/air flames.

Numerical Methodology

Numerical solutions for the one-dimensional and counterflow flames were obtained using well-established formulations and computer codes [6,9–12]. The counterflow code was further modified for more efficient convergence, more accurate calculation of the mass diffusion velocities, and the ability to obtain solutions for increasing and decreasing nozzle separation distance through continuation. Both codes were integrated to the Chemkin-II [13] and Transport [14] subroutine packages, which provide the detailed chemistry and transport information. The kinetic scheme used was a hierarchically developed C_2 mechanism [15,16], which satisfactorily predicts a wide range of oxidation properties of hydrogen, carbon monoxide, methane, ethane, ethylene, acetylene, and methanol [15,16,17–19]. For the numerical determination of the propane/air laminar flame speeds, the C_3 kinetics of Pitz and Westbrook [20] were added to the C_2 mechanism.

In addition to solving the one-dimensional unstrained laminar flames of hydrogen/air, methane/air, and propane/air mixtures, the counterflow flame was also solved for lean hydrogen/air mixtures in order to identify the influence of the separation distance on such weak flames. For the counterflow solutions, zero velocity gradient was used at the nozzle exit, and the

nozzle separation distances were 7, 14, 22, and 50 mm.

Typical numerically determined velocity profiles of the counterflow flame are given in Fig. 1 for a fixed strain rate of $K = 308 \text{ s}^{-1}$. Upon exiting the nozzle, the velocity gradually develops an increasing slope, which becomes maximum just before the minimum velocity point at which the heating starts. This maximum velocity gradient in the hydrodynamic zone is defined as the imposed strain rate, K , and the minimum velocity is a reference upstream flame speed, $S_{u,\text{ref}}$, as mentioned earlier. Typically, a numerical solution is obtained at the lowest possible strain rate K required for convergence, and then, by using continuation for the nozzle exit velocity, the dependence of $S_{u,\text{ref}}$ on K is determined.

The calculations were performed by using upwind differencing for the convective terms since convergence with the central difference has been found to be difficult in our computational facilities. Similarly, it was found in an earlier study [8] that the effect of numerical diffusivity, introduced by using upwind differencing, can be minimized by further refining the computational mesh. Thus, by increasing the number of grid points, N , the calculated flame speed decreases for the methane/air and propane/air mixtures and increases for the ultralean hydrogen/air mixtures. The reported data were obtained by plotting S_u^0 vs $1/N$ and extrapolating to $1/N = 0$, which corresponds to the limit of infinitely large N . This approach reduced the stoichiometric S_u^0 values for methane/air and propane/air mixtures by about 3 cm/s and increased the S_u^0 value for the $\phi = 0.35$ hydrogen/air mixture by 5 cm/s.

Experimental Methodology

The counterflow opposed-jet technique for the determination of laminar flame speeds and extinction strain rates is well documented [3,4,15,16,21]. In the present study, the nozzle separation distance was varied, and in agreement with our previous study [7], it was found that, by using L larger than about 14 mm, the experimental data on S_u^0 are minimally affected for methane/air and propane/air mixtures. Thus, the flame speeds for these flames were all obtained with $L = 22$ mm. For lean hydrogen/air flames, values of L equal to 7, 14, and 22 mm were used to demonstrate the influence of the finite separation distance. Efforts were made to obtain data under very low strain rates so as to minimize the stretch effect. Experiments were conducted under atmospheric pressure for the entire flammable range of methane/air and propane/air mixtures, and for lean hydrogen/air flames. For all cases, L was equal to the nozzle diameter, D , although additional measurements with the maximum $D = 22$ mm and $L = 7, 14$ mm indicated no effect on the flame speed data. This is due

to the fact that measurements were taken along the centerline, in the vicinity of which the nozzle diameter effect is not important. Furthermore, formulation of the counterflow equations realistically models our data because these equations represent an asymptotic expansion around the centerline of the system. Deviations would be expected if measurements were taken at radial distances of the same order as the nozzle radius. The uncertainty of the reported experimental data is limited by the accuracy of the LDV, which is estimated to be about ± 1 cm/s.

Results on Ultralean Hydrogen/Air Flames

The effect of the finite domain on S_u^0 was first numerically examined for the ultralean, $\phi = 0.3$ hydrogen/air mixture at $L = 7, 14$, and 22 mm. This range of the nozzle separation distance is representative of that adopted in previous experiments. We expect that the influence of L is most severe for these mixtures. Figure 1 clearly shows that the reference velocity, $S_{u,ref}$, increases with decreasing L for the same value of K . That is, since the influence of the finite flame thickness on the flow velocity increases with decreasing L [7], as the freestream flow approaches the flame, the effect of thermal expansion is more readily felt by the flow for small L . This leads to an earlier turnaround in the velocity profile and hence a higher $S_{u,ref}$, as well as a higher strain rate within the reaction zone [22]. It is further seen that the influence is especially significant for $L = 7$ mm and diminishes for larger values of L .

In Fig. 2, $S_{u,ref}$ is plotted vs the strain rate K for the three separation distances. It is seen that overall the data show a curved behavior, which becomes more prominent at the lower K values, and that $S_{u,ref}$ is uniformly higher for smaller separation distances. The dependence on L , however, diminishes with increasing L such that the differences between the results for $L = 14$ and 22 mm appear to be quite small. A linear extrapolation of the $L = 7$ mm and 22 mm data would yield flame speeds of about 15 and 10 cm/s, respectively. Such a significant difference was not observed for the previous study of methane/air flames [7], which showed only about 10% difference. Furthermore, even the 10 cm/s data for $L = 22$ mm substantially overpredict the independently calculated one-dimensional laminar flame speed of 6 cm/s.

In order to identify the observed sensitivity on the separation distance, we have evaluated the Karlovitz number experienced by the flame, defined as $Ka = K\delta/S_u^0$, where δ is a characteristic flame thickness. Since Ka represents the ratio of the characteristic residence time in the flame zone to that of the hydrodynamic zone, a smaller Ka implies a thinner flame and hence a correspondingly smaller influence of the finite separation distance. To evaluate δ , the basic laminar flame structure requires that δ be re-

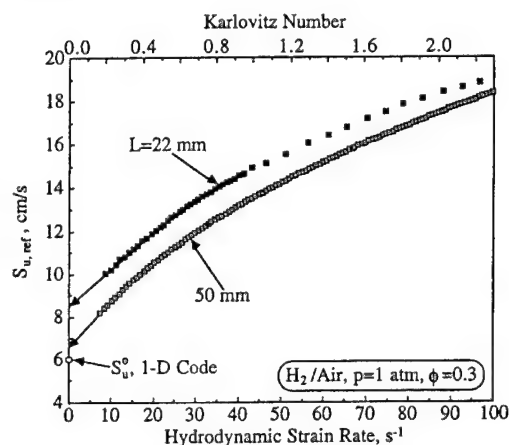


FIG. 3. Variation of the numerically determined $S_{u,ref}$ for low values of K and Karlovitz number, for $\phi = 0.3$ atmospheric hydrogen/air mixture and $L = 22$ and 50 mm.

lated to S_u^0 through $\delta(\rho_u S_u^0) = (\lambda/c_p)_u$, where ρ_u is the density of the unburned gas, λ the thermal conductivity, and c_p the specific heat, and the subscript "u" designates the unburned mixture. It represents the characteristic diffusion thickness and is well defined once S_u^0 is given. Alternatively, an empirical flame thickness can also be defined by measuring some effective thicknesses of the experimentally or numerically determined temperature profile across the flame, for example by using the tangent definition [23], which would yield a flame thickness typically larger than the diffusion thickness by a factor of about 3 and more for the present flame. Either definition can be used for the present purpose, as long as it is used consistently in the comparisons. Thus, by using the tangent method, the Karlovitz numbers for the strain rates of Fig. 2 have been evaluated and indicated. It is seen that, even for the lowest strain rate of $K = 50$ s $^{-1}$ obtainable in the experiments, the Karlovitz number is still $O(1)$. This is to be contrasted with the values of 0.1 – 0.2 (for the same range of strain rates) obtained for methane/air flames [7], for which linear extrapolation yields values close to the independently calculated laminar flame speeds. This exceptionally large Ka is due to the very small flame speeds and larger flame thicknesses for these lean mixtures.

In order to further improve the accuracy of the extrapolation, we have extended the calculations for $L = 22$ mm down to lower rates of strain such that Ka is around 0.2 , as shown in Fig. 3. It is seen that the extrapolation now places the flame speed to 8.5 cm/s, which is closer to the laminar flame speed of 6 cm/s.

An alternate approach to increase the accuracy is to further increase the separation distance. Hence, by using $L = 50$ mm and very small strain rates, Fig.

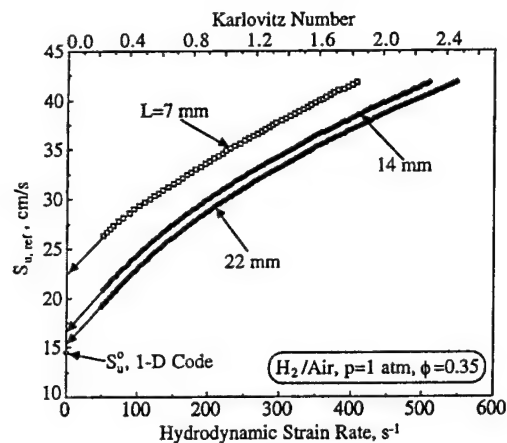


FIG. 4. Variation of the numerically determined $S_{u,ref}$ with K and Karlovitz number for $\phi = 0.35$ atmospheric hydrogen/air mixture and $L = 7, 14$, and 22 mm.

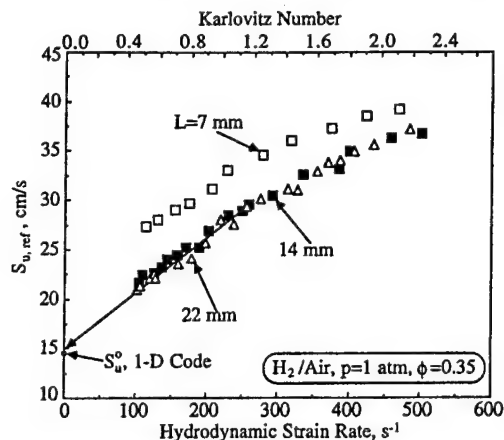


FIG. 6. Variation of the experimentally determined $S_{u,ref}$ with K and Karlovitz number for $\phi = 0.35$ atmospheric hydrogen/air mixture and $L = 7, 14$, and 22 mm.

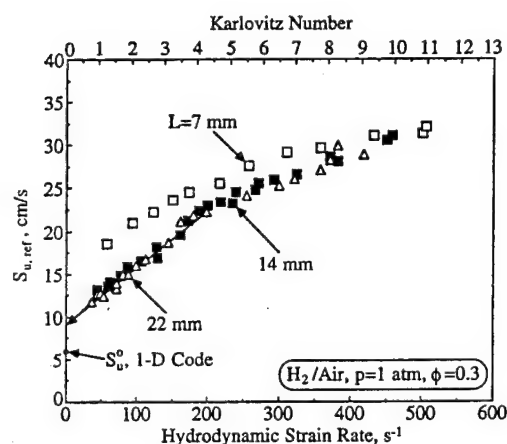


FIG. 5. Variation of the experimentally determined $S_{u,ref}$ with K and Karlovitz number for $\phi = 0.30$ atmospheric hydrogen/air mixture and $L = 7, 14$, and 22 mm.

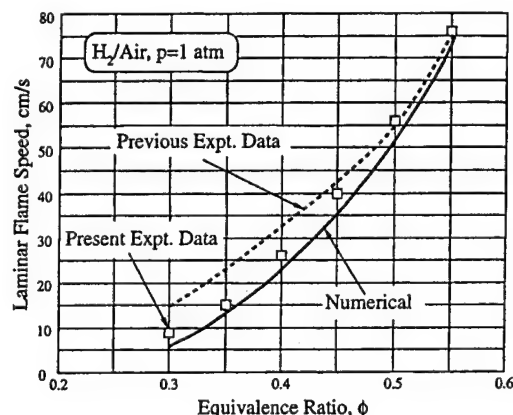


FIG. 7. Experimentally and numerically determined laminar flame speeds, S_u^0 , as a function of the equivalence ratio, ϕ , for lean atmospheric hydrogen/air mixtures.

3 shows that the extrapolated value is now 6.6 cm/s, which is within 10% of the laminar flame speed.

Figure 4 shows the corresponding case of $\phi = 0.35$. Here, even though the change in ϕ from 0.3 is slight, the calculated S_u^0 increases to 14.5 cm/s. Thus, the Karlovitz number is reduced for the same physical strain rate. A linear extrapolation with $L = 22$ mm and $K = 50$ s $^{-1}$ in this case yields 15.5 cm/s, which is close to the laminar flame speed.

Figures 5 and 6 show the experimental results for the $\phi = 0.3$ and $\phi = 0.35$ hydrogen/air mixtures, respectively, with $L = 7, 14$, and 22 mm. The minimum strain rates that could be established for $\phi = 0.3$ and 0.35 , with $L = 22$ mm, without inducing flashback in our system were about 50 and 100 s $^{-1}$,

respectively. It is seen that, since the minimum Ka is about 1 for the $\phi = 0.3$ flames, accuracy of the linear extrapolation is not assured. Indeed, extrapolation yields 9 cm/s, which still substantially overpredicts the calculated S_u^0 of 6 cm/s. This overprediction was anticipated from the numerical solution of Fig. 2. However, for the $\phi = 0.35$ flame, the minimum Karlovitz number is about 0.4 . Consequently, the extrapolation is quite accurate, again in agreement with the numerical results.

Figure 7 compares the experimentally measured laminar flame speeds with the independently computed ones over the range of $\phi = 0.3$ – 0.55 . It is seen that, apart from the $\phi = 0.3$ value, which shows a relatively higher experimental value, comparisons for all the other cases are very close in terms of the rel-

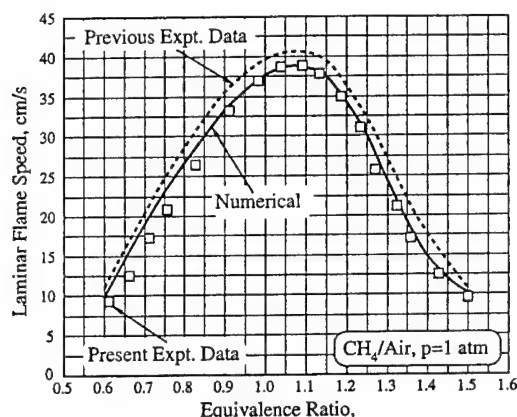


FIG. 8. Experimentally and numerically determined laminar flame speeds, S_u° , as a function of the equivalence ratio, ϕ , for atmospheric methane/air mixtures.

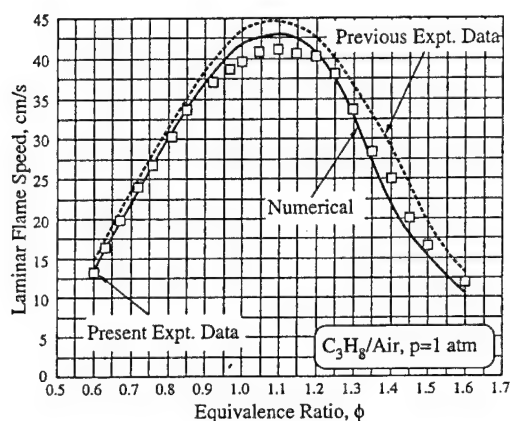


FIG. 9. Experimentally and numerically determined laminar flame speeds, S_u° , as a function of the equivalence ratio, ϕ , for atmospheric propane/air mixtures.

active extent of deviations. We have also plotted the experimental values determined previously [8] with larger strain rates. It is seen that these values are substantially higher than the calculated values. Thus, the present result provides a satisfactory resolution of this previously observed discrepancy.

Results on Methane/Air and Propane/Air Flames

Figures 8 and 9, respectively, compare the present experimental data, all obtained with a nozzle separation distance of 22 mm, with the previous data [4,15,16,21], obtained mostly with $L = 7$ and 14 mm, for methane/air and propane/air mixtures. The comparison shows that the present experimental flame

speeds are indeed lower than the previous values, particularly for rich propane/air mixtures. The differences, however, are much smaller than those for the ultralean hydrogen/air flames. This is reasonable based on considerations of the Karlovitz number for the experimental conditions. In addition, we have also compared the present experimental data with the calculated data by using the one-dimensional flame code. The comparison also shows close agreement, especially for the methane/air flames.

It may be noted that, in our previous studies for the methane and propane flames [4,15,16,21] using smaller separation distances, reasonably good agreements between computed and experimental values were also obtained. Such an agreement is a consequence of the higher experimental flame speeds due to the small L used, and the higher computed flame speeds based on previously accepted kinetic data. Agreement is again found in the present study because, while the experimental flame speeds are now reduced, the computed values obtained by using recent kinetic data and improved numerical accuracy are also reduced. The important points to note are that the computation and experiments were always independently conducted, that the differences between the previous and present values are actually quite small, that the present results are consequences of improvements in chemical kinetics and the experimental and numerical techniques, and that the present agreement also includes the rich propane/air flames.

Concluding Remarks

In this study, we have provided further insight into the accuracy of the linear extrapolation associated with the determination of laminar flame speeds by using the counterflow twin-flame technique, and identified approaches toward minimizing the inaccuracies. Specifically, it is shown that the accuracy of the linear extrapolation increases with decreasing Karlovitz number, that the Karlovitz number needs to be reduced to $O(0.1)$, based on the tangent definition of the flame thickness, in order for the linear extrapolation to be accurate, and that the accuracy can be improved by either reducing the strain rate or increasing the nozzle separation distance. Perhaps the most significant consequence of the above considerations is that our redetermined laminar flame speeds of ultralean hydrogen/air mixtures now agree well with the independently calculated values, thereby satisfactorily resolving the previously observed discrepancy of the much higher values of the experimental data. The redetermined values of the methane/air flames also show closer agreement with the calculated values obtained with the recent kinetic data of methane oxidation.

Under most common situations, the counterflow

technique with linear extrapolation is expected to yield flame speeds of sufficient accuracy, provided the lowest strain rate is not too high while the nozzle separation distance is also not too small. However, since strain rate reduction can be limited by flashbacks while increasing the nozzle separation distance, and hence nozzle diameter can also be complicated by buoyancy-induced flame instability and other experimental difficulties, the accuracy of this technique may decrease for the determination of the laminar flame speeds of very weak flames. We are currently developing nonlinear extrapolation approaches to circumvent such limitations.

Acknowledgments

CMV and FNE were supported by the National Science Foundation under Grant No. NSF CTS-9211844. CKL was supported by the Air Force Office of Scientific Research and the Office of Naval Research.

REFERENCES

- Andrews, G. E., and Bradley, D. *Combust. Flame* 18:133-153 (1972).
- Markstein, G., *Nonsteady Flame Propagation*, MacMillan, New York, 1964, pp. 5-73.
- Wu, C. K., and Law, C. K., *Twentieth Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1984, pp. 1941-1949.
- Zhu, D. L., Egolfopoulos, F. N., and Law, C. K., *Twenty-Second Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1988, pp. 1537-1545.
- Tien, J. H., and Matalon, M., *Combust. Flame* 84:238-248 (1991).
- Dixon-Lewis, G., *Twenty-Third Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1990, pp. 305-324.
- Chao, B. H., Egolfopoulos, F. N., and Law, C. K., "Structure and Propagation of Counterflow Premixed Flames in a Finite Domain," manuscript in preparation.
- Egolfopoulos, F. N., and Law, C. K., *Twenty-Third Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1990, pp. 333-340.
- Kee, R. J., Grcar, J. F., Smooke, M. D., and Miller, J. A., *A Fortran Program for Modeling Steady Laminar One-Dimensional Premixed Flames*, Sandia Report SAND85-8240, 1985.
- Miller, J. A., Kee, R. J., Smooke, M. D., and Grcar, J. F., "The Computation of the Structure and Extinction Limit of a Methane-Air Stagnation Point Diffusion Flame," Spring Meeting of the Western States Section of The Combustion Institute, Paper WSS/CI 84-10, 1984.
- Kee, R. J., Miller, J. A., Evans, G. H., and Dixon-Lewis, G., *Twenty-Second Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1988, pp. 1479-1494.
- Kee, R. J., personal communications.
- Kee, R. J., Rupley, F. M., and Miller, J. A., *Chemkin-II: A Fortran Chemical Kinetics Package for the Analysis of Gas-Phase Chemical Kinetics*, Sandia Report SAND89-8009, 1989.
- Kee, R. J., Warnatz, J., and Miller, J. A., *A FORTRAN Computer Code Package for the Evaluation of Gas-Phase Viscosities, Conductivities, and Diffusion Coefficients*, Sandia Report SAND83-8209, 1983.
- Egolfopoulos, F. N., Cho, P., and Law, C. K., *Combust. Flame* 76:375-391 (1989).
- Egolfopoulos, F. N., Zhu, D. L., and Law, C. K., *Twenty-Third Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1990, pp. 471-478.
- Egolfopoulos, F. N., Du, D. X., and Law, C. K., *Combust. Sci. Technol.* 83:33-75 (1992).
- Vagelopoulos, C. M., and Egolfopoulos, F. N., "Laminar Flame Speeds and Extinction Strain Rates of Mixtures of Carbon Monoxide with Hydrogen, Methane, and Air," *Twenty-Fifth Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1994, pp. 1317-1323.
- Egolfopoulos, F. N., and Law, C. K., "Further Considerations on the Methanol Oxidation Kinetics," manuscript in preparation.
- Pitz, W. A., and Westbrook, C. K., *Combust. Flame* 63:113-133 (1986).
- Law, C. K., Zhu, D. L., and Yu, G., *Twenty-First Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1986, p. 1419.
- Egolfopoulos, F. N., "Geometric and Radiation Effects on Steady and Unsteady Strained Laminar Flames," *Twenty-Fifth Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1994, pp. 1375-1381.
- Tsatsaronis, G., *Combust. Flame* 33:217-239 (1978).



OPPORTUNITIES AND CHALLENGES OF COMBUSTION IN MICROGRAVITY

C. K. LAW* and G. M. FAETH†‡

*Department of Aerospace and Mechanical Engineering, Princeton University, Princeton, NJ 08544-5263, U.S.A.
 †Department of Aerospace Engineering, The University of Michigan, Ann Arbor, MI 48109-2118, U.S.A.

Received 21 February 1994

Abstract—This review considers the opportunities for enhanced fundamental combustion understanding from experiments where effects of buoyancy are eliminated, and the new challenges of fire safety considerations in nonbuoyant (spacecraft) environments. The following specific microgravity combustion phenomena are considered: stretched flames, flamefront instabilities, flammability limits and near-limit phenomena of gaseous premixed flames; structure, stability and soot processes in gaseous nonpremixed flames; flame propagation, smoldering and materials synthesis in heterogeneous premixed flames; flame spread, gasification and combustion in heterogeneous nonpremixed flames; flame-inhibiting atmospheres, fire detection and extinguishment in spacecraft environments; and ground-based (drop tower and aircraft), sounding rocket and space-based (shuttle, space station) microgravity combustion research facilities that are either available or anticipated.

The findings of the review highlight how buoyancy has impeded the rational development of combustion science, precluding observations of fundamental one-dimensional configurations, low Reynolds number flows and other limiting conditions that have been invaluable for developing understanding in other areas of science. Thus, experiments at microgravity provide an opportunity to finally merge theories and experiments for classical problems in order to advance the fundamental understanding of combustion phenomena. Additionally, combustion processes have been shown to be very different at normal gravity and microgravity so that improved understanding of combustion at microgravity is needed in order to address fire and explosion safety considerations for spacecraft.

CONTENTS

Notation	66
1. Introduction	66
2. Intrusion of Buoyancy	67
3. Gaseous Premixed Flames	69
3.1. Introduction	69
3.2. Stretch	70
3.3. Flamefront instability	74
3.4. Flammability limits	77
3.5. Near-limit phenomena	80
3.5.1. Self-extinguishing flames	80
3.5.2. Stationary spherical flames	82
4. Gaseous Nonpremixed Flames	83
4.1. Introduction	83
4.2. Flame structure	84
4.3. Flame stability	87
4.4. Soot processes	88
5. Heterogeneous Premixed Flames	92
5.1. Introduction	92
5.2. Flame propagation in suspensions	93
5.3. Smoldering	94
5.4. Materials synthesis	95
6. Heterogeneous Nonpremixed Flames	95
6.1. Introduction	95
6.2. Flame spreading	95
6.3. Droplets	98
7. Spacecraft Fire Safety	103
7.1. Introduction	103
7.2. Flame-inhibiting atmospheres	104
7.3. Fire detection and extinguishment	105
8. Microgravity Facilities	106
8.1. Introduction	106
8.2. Drop Towers	106

‡Author to whom correspondence should be addressed.

8.3. Aircraft	107
8.4. Sounding rockets	108
8.5. Spacecraft	108
9. Concluding Remarks	109
Acknowledgements	109
References	109

NOTATION

c	molar concentration	μ_F	characteristic forced convection velocity
C_p	specific heat at constant pressure	u_N	characteristic natural convection velocity
d	burner exit diameter	v_f	flame spreading velocity
d_s	drop diameter	\dot{w}	reaction rate per unit volume
d_f	flame diameter	x	streamwise distance
D	mass diffusivity	y	cross stream distance
e	base of the natural system of logarithms	Y_O	oxidant mass fraction
f_s	soot volume fraction	Y_P	combustion product mass fraction
g	acceleration of gravity	Y_R	reactant mass fraction
Gr	Grashof number, Eq. (1)	z	streamwise distance
K	flame stretch	α	thermal diffusivity
Ka	Karlovitz number	δA	increment of flame surface area
l_K	Kolmogorov length scale of turbulence	δ_L	flame thickness
L	characteristic length, flame length	$\Delta\rho$	density difference
Le	Lewis number	ε	rate of dissipation of turbulence kinetic energy
L_M	Markstein length	λ	thermal conductivity
Mk	Markstein number	Λ	integral scale of turbulence
ng	normal gravity	μg	microgravity
Pe	Peclet number for liquid mass diffusion	ν	kinematic viscosity; stoichiometry parameter, $Y_{O\infty}/\sigma$
Pr	Prandtl number, α/ν	ρ	density
p	pressure	σ	stoichiometric mixture ratio
r	radial distance	τ_K	Kolmogorov timescale of turbulence
r_f	flame radius	ϕ	fuel-equivalence ratio
Re	Reynolds number, Eq. (5); burner exit Reynolds number, du_o/v_o	τ_L	characteristic time of a laminar flame
Re_Λ	turbulence Reynolds number, $\bar{u}'\Lambda/\nu_u$	X_F	transport parameter, $\lambda_F/(cD_F)$
R_f	flame radius		
Ri	Richardson number, Eq. (3)		
Sc	Schmidt number, D/ν		
S_L	laminar burning velocity		
t	time		
T	temperature		
T_f	flame temperature		
T_{ad}	adiabatic flame temperature		
u	streamwise velocity		
U	streamwise velocity		

Subscripts

b	combustion products or burned gas property
F	fuel property
u	unreacted gas property
o	burner exit condition; initial condition
∞	ambient condition; negligible stretch condition

Superscripts

$(-)$	mean value
$(-)'$	rms fluctuating value

1. INTRODUCTION

The availability of improved microgravity (μg) facilities and increased manned space activities offer unprecedented opportunities and challenges to combustion science and technology: opportunities to study many combustion phenomena whose fundamental understanding has been handicapped thus far by the buoyant flows that accompany flame processes on Earth, and challenges to develop rational preventive guidelines and control strategies for fire and explosion hazards for spacecraft.¹⁻⁵ The objectives of this review are to discuss progress toward both exploiting the opportunities and meeting the challenges of combustion at μg conditions, concluding with a description of available and anticipated facilities for μg combustion research.

Both terrestrial and space applications provide motivation for advancing fundamental understand-

ing of combustion science. In particular, we are currently confronted with a long list of either unresolved or emerging combustion problems that have strong economic, social, political and defense-related relevance on Earth. Examples are energy conservation and utilization, air pollution, surface-based transportation, aircraft and spacecraft propulsion, municipal and hazardous waste incineration, materials processing and synthesis, and atmospheric change and global warming. Furthermore, uncontrolled fires and explosions continue to cause loss of life and property—problems that will only become worse as our population and the application of novel technologies increase. Finally, our inadequate understanding of fire and explosion phenomena on Earth is exacerbated by the novel nonbuoyant environment of spacecraft; clearly, combustion research must merit high priority in the space program in order to avoid combustion-related tragedies in the future.

Several reviews of aspects of μg combustion science have appeared recently.¹⁻⁵ References 1 and 2 are overviews of the U.S. Microgravity Combustion Science Program up to 1989 and 1992, respectively. Law³ provides a review emphasizing the advantages of combustion experiments at μg conditions and highlighting the need for studies of spacecraft fire safety due to documented effects of gravity on the properties of flames. Faeth⁴ reviews studies of laminar premixed and nonpremixed gas flames at μg , updating aspects of Refs 1 and 2. Finally, Sacksteder⁵ describes available ground-based and spacecraft facilities for μg combustion research and provides a summary of current experiments using these facilities. The present paper synthesizes and updates these reviews up to 1993, emphasizing accomplishments and issues for both homogeneous and heterogeneous flames. The discussion is limited to fundamental laminar flame processes, basic to both laminar and turbulent flames, because practical turbulent flames will remain an empirical aspect of combustion science for some time to come due to computational and experimental limitations.⁶

The paper begins with a discussion of the intrusion of buoyancy during normal gravity (ng) measurements of flame properties, in order to highlight potential research opportunities using μg environments. Completed and active studies of the following combustion phenomena at μg are then considered in turn: stretch, flamefront instabilities, flammability limits and near-limit phenomena of gaseous premixed flames; structure, stability and soot processes in gaseous nonpremixed flames; flame propagation, smoldering and materials synthesis in heterogeneous premixed flames; flame spread, gasification and combustion in heterogeneous nonpremixed flames; and flame-inhibiting atmospheres, fire detection and fire extinguishment in spacecraft environments. The paper concludes with a description of available ground-based (drop tower and aircraft), sounding rocket and space-based (shuttle, space station) μg combustion research facilities and a summary of recommendations for future μg combustion research based on the needs of combustion-related technologies and the capabilities of available μg combustion research facilities.

2. INTRUSION OF BUOYANCY

The intrusion of gravitational forces is a greater impediment to combustion studies than most other areas of science. By its nature, combustion involves chemical energy releases which typically cause the temperature of reactive mixtures to increase from an unreacted ambient state of roughly 300 K to totally reacted states at 2000–3000 K. These large temperature differences yield corresponding density differences, which in the presence of gravity invariably cause buoyant motions that vastly complicate both

the execution and interpretation of measurements. Buoyant motion also prevents some fundamental phenomena—like most laminar one-dimensional premixed and diffusion flames, low Reynolds number heterogeneous flames, flame spread in dispersed heterogeneous media, etc.—from being observed at all. Perversely, the problems of buoyancy are greatest for fundamental laboratory experiments where accurate temporal and spatial resolution are important factors.

Simple phenomenological considerations help to quantify the limitations caused by buoyancy during fundamental flame experiments and provide insight concerning the focus of current μg combustion research. Molecular transport (as manifested by the diffusion of mass, momentum and thermal energy) and forced motion are the relevant collateral properties to chemical energy release for most combustion phenomena; in contrast, buoyant motion generally is an unwanted intrusion. The relative importance of the collateral properties and buoyancy can be represented by two dimensionless parameters: the ratio of buoyant to molecular transport, called the Grashof number, and the ratio of buoyant to forced convective transport, called the Richardson number.⁷

The Grashof number, Gr , is defined as follows:

$$Gr = (\Delta\rho/\rho)gL^3/\nu^2 \quad (1)$$

where $\Delta\rho$ and ρ are the characteristic density difference and mean density of the process, respectively, g is the acceleration of gravity, L is a characteristic length scale of the process, and ν is a mean kinematic viscosity. Noting that $\Delta\rho/\rho \approx 1$ for flames, because the density of the reactants is generally large in comparison with the density of the combustion products, and that $Gr < O(10^{-1})$ for effects of buoyancy to be small,⁷ we find that

$$L < O(100 \mu m) \quad (2)$$

for typical values of ν (ca. 10 mm²/s) in atmospheric pressure flames. Unfortunately, it is not possible to resolve experiments on such scales using either existing or anticipated combustion apparatus and instrumentation. Experiments at subatmospheric pressures can increase allowable flame sizes before buoyancy intrudes, and this has been exploited by some workers, for example, Law *et al.*⁸ for studies of droplet burning in quiescent environments. However, reduced rates of chemical reaction at low pressures, even with oxygen enrichment, allow only small increases in scales ($L \sim p^{-2/3}$ from Eq. (1), where p denotes pressure) before flame extinction eventually occurs. Furthermore, subatmospheric tests generally have reduced relevance because most applications involve pressures equal to or greater than atmospheric pressure, while extrapolation of low-pressure results to the range of interest is uncertain due to complex effects of pressure on the large number of individual reaction steps normally involved in combustion chemistry.

Experiments in the presence of finite flow velocities offer a way of circumventing the Grashof number limitations. The Richardson number, Ri , is a measure of the relative importance of buoyancy and a characteristic velocity associated with the flame, such as the laminar burning velocity of premixed flames, S_L , or the forced convection velocity of non-premixed flames, u_F . The physical basis for this parameter can be seen by finding the characteristic natural convection velocity, $u_N = (\Delta\rho g L / \rho)^{1/2}$, when a low-density gas rises a distance L .⁷ This characteristic natural convection velocity is based on a balance between buoyant and inertial forces; the effects of viscous forces will be considered subsequently. Forming the ratio of the square of these velocities we have:

$$Ri = (\Delta\rho/\rho)gL/(S_L \text{ or } u_F)^2. \quad (3)$$

In the following, we will consider the implications of Eq. (3) for premixed and nonpremixed flames in turn.

Reasonable flame dimensions to provide adequate spatial resolution for measurements within premixed laminar flames are in the range 10–100 mm, while $Ri < O(10^{-1})$ for effects of buoyancy to be small. We then find that

$$S_L > 0(1 \text{ m/s}) \quad (4)$$

is required to avoid significant effects of buoyancy. This regime is comparable with maximum laminar burning velocities. However, it is not possible to study the region of small laminar burning velocities near flammability limits on Earth (typically a few centimeters per second as discussed later) without the intrusion of buoyancy. Additionally, studies of flamefront instabilities, which tend to be most important near flammability limits and for turbulent flames, are constrained by the criterion of Eq. (4) because effects of instabilities tend to be weak near high burning velocity conditions.^{9–12} We will return to this issue later.

The convective limitation due to buoyancy also affects nonpremixed flames. Effects of diffusional transport of heat and mass are central issues for nonpremixed (diffusion) flames. We can conveniently interpret these effects in terms of momentum transport, however, because ratios of heat, mass and momentum diffusivities are nearly unity for gaseous environments of interest to combustion, that is, the Prandtl, Pr , and Schmidt, Sc , numbers are nearly unity for gases. It is then convenient to interpret transport effects through the kinematic viscosity by defining a characteristic Reynolds number of these flames, Re , as follows:

$$Re = Lu_F/\nu = (Gr/Ri)^{1/2} \quad (5)$$

where Gr and Ri have been introduced from Eqs (1) and (3). Noting that $Gr = O(10^3)$ for a flame length of 10 mm (which provides reasonable spatial resolution for measurements) from Eq. (1), while

$Ri < O(10^{-1})$ for forced motion to dominate buoyancy,⁷ we have as a consequence

$$Re > 10^2. \quad (6)$$

This implies that the Stokes flow regime ($Re < 1$), which is a natural limit that has been invaluable for understanding fluid mechanics, cannot be reached for flame studies on Earth without the intrusion of buoyancy. Thus, buoyant, nonpremixed flames are a common experimental configuration, for lack of an alternative, although buoyancy causes complications and introduces phenomena that have little relevance to most studies. Furthermore, the large buoyant velocities cause flames to develop thin boundary layer-like structures typical of large Re conditions that can significantly limit the spatial resolution of measurements, in comparison with low Re conditions achievable at μg .

The effect of buoyancy is so ubiquitous that we generally do not appreciate the enormous negative impact that it has had on the rational development of combustion science. For example, aside from limited exploratory work at μg conditions, we have never observed the most fundamental processes of combustion without substantial disturbances of buoyancy—precluding simple one-dimensional configurations, low Reynolds number flows and other limiting conditions that have been invaluable for developing understanding in other areas of science. This prevents the rational merging of theory, where buoyancy is frequently of little interest, and experiments, which are always contaminated by effects of buoyancy at ng conditions.

Turbulent flames, one of the most important unresolved problems of combustion science, provide graphic examples of the impediment that buoyancy causes to the parallel development of theory and experiment. Three-dimensional time-dependent numerical simulations of turbulent flames offer a logical way to study some of the phenomena of turbulence; however, due to computer limitations, such calculations can only consider low-speed flows having a relatively limited range of length scales, that is, low Reynolds number flows.⁶ Unfortunately, such conditions cannot be duplicated in the laboratory at ng because buoyancy immediately accelerates the flow from any initial low-speed condition, resulting in high-speed flows with a large range of length scales, that is, high Reynolds number flows. The inability to measure combustion properties in low-speed two-phase flows, because particles and drops settle at ng conditions, is another obvious example of the experimental limitations caused by buoyancy. In these circumstances, theory and experiment tend to go their own way—to the detriment of both. Furthermore, even the most optimistic estimates of rates of computer development imply no merger of theoretical capabilities and experimental conditions for simulations of turbulence and practical multiphase flows for the foreseeable future.⁶ These are only examples, and

similar buoyancy-induced gaps between theory and experiment exist in virtually every area of combustion science. With no massive breakthrough in computer technology in the offing, rapid expansion of combustion experiments at μg conditions offers the most promising approach toward resolving this theoretical/experimental dichotomy of combustion science.

The same features that make the μg environment attractive for fundamental combustion experiments introduce hazards of fires and explosions that have no counterpart on Earth. Thus far, activities in space have been limited and have only involved carefully selected and highly trained individuals. As the space environment is exploited, however, a larger range of activities and individuals will be involved—vastly increasing the potential for unwanted fires and explosions in a highly public arena. The main concern is that virtually all existing information concerning design procedures to control fires and explosions is based on experience with ng environments. Of necessity, current qualification procedures for materials used in space involve tests at ng , justified by rather limited measurements from μg environments.⁵ Since we know that combustion processes are very different at ng and μg conditions, there is little basis for confidence that this practice is sufficient.

Thus, the Earth's gravity has impeded combustion science, somewhat analogously to the way that the Earth's atmosphere has impeded optical astronomy; additionally, there is virtually no technology base for fire and explosion safety considerations for spacecraft. To circumvent these difficulties, a variety of facilities for reduced gravity combustion research have been developed, including drop towers providing 1–10 s at 10^{-4} – $10^{-6} g$, aircraft-based facilities providing 5–15 s at 10^{-1} – $10^{-3} g$, sounding rockets providing up to 200–900 s at $10^{-4} g$ and spacecraft facilities providing times of 10^3 – 10^4 s at $10^{-5} g$.⁵ In the following sections we shall discuss some combustion phenomena that are strongly influenced by the presence of buoyancy, the progress using available μg facilities to study fundamental combustion processes, and some issues that remain to be resolved.

3. GASEOUS PREMIXED FLAMES

3.1. Introduction

The large reaction rates per unit volume required for efficient and compact combustion in flowing premixed gases, as well as the large velocities caused by buoyancy for flames in still gases, imply that most practical gaseous premixed flames are turbulent. Thus, an important issue that must be addressed is the relevance of laminar premixed flame studies to applications involving turbulent premixed flames. This issue has been resolved by the development of the laminar flamelet concept in recent years and the finding that most practical combustion systems are

in the laminar flamelet regime.^{13,14} The laminar flamelet concept implies that premixed turbulent flames can be considered to consist of quasi-steady premixed laminar flamefronts, that is, flame surfaces that are distorted by turbulence when certain conditions are met as discussed in the following.^{13–15} First, the laminar flamelet concept applies when the smallest scales of turbulence (which can be represented by the Kolmogorov microscale, l_K , because scales relevant to mass, momentum and energy transport are comparable in gases, as noted earlier) are larger than the characteristic flame thickness, of δ_L , that is, when $l_K > \delta_L$. This criterion can be expressed in terms of the velocity fluctuations of turbulence in the unburned gas relative to the laminar burning velocity, \bar{u}'/S_L , and the turbulence Reynolds number of the unburned gas, $\bar{u}'\Lambda/\nu_u$, where Λ is the integral scale of the turbulence, as follows:¹³

$$\bar{u}'/S_L < Re_\Lambda^{1/2}, \text{ laminar flamelet regime.} \quad (7)$$

Additionally, the quasisteady requirement can be met when the Kolmogorov microtime scale of the unburned gas, $\tau_K = (\nu_u/\epsilon)^{1/2}$, where ϵ is the rate of dissipation of turbulence kinetic energy, is greater than the characteristic residence time of the flame, $\tau_L = \delta_L/S_L$, or $\tau_K > \tau_L$. It is easy to show, using an argument similar to Bray,¹³ that the quasisteady approximation also requires the conditions specified by Eq. (7). Thus, the laminar flamelet regime involves distortion of the laminar flame sheet at scales that are larger than the thickness of the flame and at rates that are slow in comparison with the response capabilities of the flame. When these conditions are satisfied, the properties of laminar flames are relevant to turbulent flames. Furthermore, the flame thickness requirement expressed by Eq. (7) is not very restrictive; for example, even the intense turbulent premixed flames found within automotive spark-ignition engines are largely within the laminar flamelet regime.¹⁶ Thus, the effects of turbulence for a significant range of practical conditions can be viewed in terms of enhanced or retarded (quenched) laminar flamelet propagation due to turbulence-induced flame distortion. These distortions involve wrinkling of the propagating laminar flamefront by the nonuniformities of the turbulent flow field, which introduces variations in the curvature of the flamefront, as well as regions where the flamefront is compressed or expanded (strained) along its surface. These dynamical effects of turbulence on thin laminar flamefronts are reviewed by Peters¹⁴ and Law,¹⁵ where it is shown that the combined effects of curvature and strain on a propagating thin laminar flamefront can be conveniently represented by the flame stretch. Flame stretch, which was introduced by Karlovitz *et al.*,¹⁷ is defined as the local fractional rate of increase of flame surface area. Thus, effects of distortion of laminar premixed flames, characterized as effects of flame stretch, are receiving significant attention, including work at μg conditions.

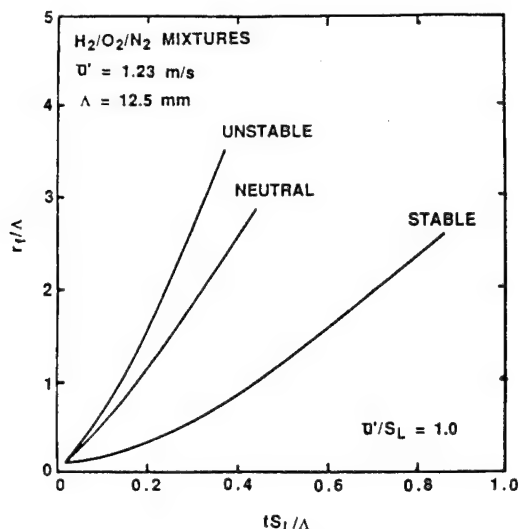


FIG. 1. Variation of the properties of freely propagating turbulent premixed flames in $H_2/O_2/N_2$ mixtures due to effects of preferential diffusion. From Tseng and Faeth.¹⁸

Another aspect of the relevance of properties of laminar flamefronts to turbulent premixed flames involves potential effects of laminar flamefront instabilities. As discussed subsequently, the interaction between flow nonuniformities in a turbulent flow field and a laminar flamefront generally is not passive, with effects of flamefront instability acting to either enhance or retard the turbulent distortion of the flame surface for unstable and stable conditions, respectively. There are three main mechanisms of laminar flamefront instability, as reviewed by Clavin¹⁸ and these will be discussed in more detail subsequently. These three mechanisms include: (i) hydrodynamic instability, which is caused by acceleration of low-density combustion products toward a high-density reactant mixture; (ii) preferential-diffusion instability, which is caused by different heat and mass transport rate response to effects of flame stretch; and (iii) Rayleigh–Taylor instabilities, which are caused by accelerations, mainly due to gravity, normal to a density discontinuity such as a flame. All three mechanisms are present to some extent for laminar flamefronts at ng, however, the preferential-diffusion mechanism appears to be most important for premixed turbulent flames.

Recent observations for a variety of turbulent premixed flames suggest that effects of preferential diffusion, analogous to those associated with preferential-diffusion instability of laminar flamefronts, generally affect the properties of turbulent premixed flames. For example, Clavin and coworkers¹⁸ have found that flamefront stability phenomena, due to effects of preferential diffusion, influence turbulent flames having large length scales and low turbulence intensity, with unstable and stable conditions, respectively, causing chaotic enhancement and damping of flame surface distortion by turbulence.

While these findings were for weak turbulence, recent experimental work supports similar behavior for highly turbulent, rim-stabilized and freely-propagating flames.^{9–12,19} An example of this behavior is illustrated in Fig. 1, which is a plot of mean flame position as a function of time for freely propagating flames in $H_2/O_2/N_2$ mixtures within the laminar flamelet regime at $Re_\Delta = 1200$ and $\bar{u}'/S_L = 1.19$. Results are illustrated for three flames having nearly identical laminar burning velocities and unburned gas turbulence properties, which involve unstable, nearly-neutral and stable preferential-diffusion stability conditions, respectively. Barring effects of preferential-diffusion instability, these flames should have identical properties because test conditions were remote from quenching limits; instead, the unstable (stable) flame propagates substantially faster (slower) than the neutrally-stable flame. This behavior clearly shows that effects of preferential diffusion are important even for highly turbulent conditions. Although effects of preferential diffusion in turbulent flames are just beginning to receive attention in the literature, they are important for most practical applications. For example, premixed flames in spark ignition automotive engines and aircraft propulsion systems all are within the strongly stable preferential-diffusion regime where distortion of the flame surface by turbulence is damped. Coupled with the laminar flamelet concept, this has motivated new interest in the classical problem of laminar flamefront instability.

The preceding discussion suggests that laminar premixed flames are relevant to turbulent premixed flames and has highlighted the importance of stretch, flamefront instability and limits (quenching) for practical flames. Studies of these topics at μg will be considered in the following, concluding with some near-limit phenomena—self-extinguishing flames (SEF) and stationary spherical flames (SSF)—that appear to be unique to μg conditions.

3.2. Stretch

Effects of flame stretch, K , are important for understanding flamefront stability and aspects of limits; they will, therefore, be discussed in this section. As noted earlier, flame stretch collectively represents the effects of flow nonuniformity, flame curvature and flame motion. It is defined in the context of a laminar flame surface element of incremental area, δA , as follows:¹⁷

$$K \equiv d \ln (\delta A) / dt \quad (8)$$

where changes of δA are observed in a Lagrangian reference frame that moves with the flame surface, that is, the boundary of δA moves in a tangent to the flame surface at the local tangential velocity of the gas, and normal to the flame surface at the sum of the local normal velocity of the gas and the local

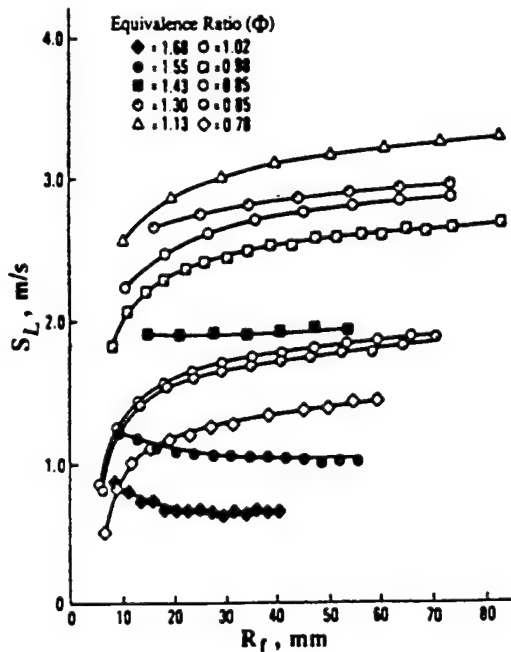


FIG. 2. Variation of laminar burning velocity with flame radius for positively-stretched propane-air flames. Data from Palm-Leis and Strehlow²⁷ as replotted by Law.¹⁵

laminar burning velocity of the flame.^{14,15} Dynamic effects due to stretch and the extent of preferential diffusion alter the relative rates of transport of gas species and thermal energy within the preheat zone of the flame structure, consequently modifying the temperature, composition and reaction rates within the reaction zone. The corresponding modification of laminar burning velocities yields the response of the flame to stretch.

Simplified theories of stretched laminar flames^{15,18-23} frequently adopt a global one-step reaction whose rate is controlled by the stoichiometrically-deficient reactant: the fuel for fuel-lean conditions or the oxidant for fuel-rich conditions. Preferential diffusion of heat and mass is allowed by considering a nonunity Lewis number, Le , which is interpreted to be the ratio of the thermal diffusivity of the bulk mixture to the mass diffusivity of the stoichiometrically-deficient reactant. For near-stoichiometric mixtures an effective Le can also be identified.²⁴ For these approximations, the flame temperature, T_f , in the linearized limit of $Ka(1 - Le)/Le \ll 1$, is related to the adiabatic flame temperature, T_{ad} , as follows:¹⁵

$$T_f T_{ad} = 1 + Ka_\infty(1 - Le)/Le. \quad (9)$$

The parameter Ka in Eq. (9), is the Karlovitz number which is the stretch of the flame surface normalized by the characteristic residence time of the flame:

$$Ka = K/\tau_L = KS_L/\delta_L \quad (10)$$

and the subscript ∞ designates conditions near the unstretched state of $K = 0$. The use of Ka_∞ instead

of Ka in Eq. (9) indicates the linearized nature of the solution. Equation (9) implies increased flame temperatures ($T_f > T_{ad}$) for either $Ka_\infty > 0$ and $Le < 1$ or $Ka_\infty < 0$ and $Le > 1$, and reduced flame temperatures ($T_f < T_{ad}$) otherwise. This result has been extensively verified experimentally¹⁵ for both counterflow and Bunsen flames, for which the stretch effect is manifested through flow straining and flame curvature, respectively.

Modification of the flame temperature through stretch implies a corresponding modification of the reaction rate, \dot{w} , and hence the laminar burning velocities, flammability limits and quenching conditions. In particular, phenomenological analysis shows that $S_L \sim \dot{w}^{1/2}$, where \dot{w} increases with increasing temperature in the Arrhenius manner (see Ref. 23). This behavior is illustrated by measurements of laminar burning velocities for free spherical flames propagating radially outward from a centrally located spark source in an initially motionless combustible mixture. If the burned gas has negligible motion, if the flame is thin in comparison with its radius, $\delta_L \ll r_f$, and if the rate of change of flame thickness with flame radius is small, $d\delta_L/dr_f \ll 1$, which usually is the case, Eq. (8) yields:^{15,25,26}

$$K = (2/r_f) dr_f/dt \quad (11)$$

where r_f is the radius of the flame surface. Furthermore, a simple mass balance between the rate of consumption of the unburned mixture ($\sim S_L \rho_u$) and the rate of growth of the burned mixture ($\sim \rho_b dr_f/dt$) yields:

$$S_L = (\rho_b/\rho_u) dr_f/dt. \quad (12)$$

Then, eliminating dr_f/dt between Eqs (11) and (12), we have:

$$K = 2(S_L/r_f)(\rho_u/\rho_b). \quad (13)$$

For the outward propagation of a spherical flame, K is positive and decreases monotonically toward zero as r_f increases. Then Eqs (9) and (10) imply that $T_f > T_{ad}$ for $Le < 1$, and vice versa for $Le > 1$.

While the transition Le for T_f is unity, the transition Le for the laminar burning velocity deviates from this value.¹⁵ Thus, flame curvature and flow nonuniformity still have an influence on the laminar burning velocity when $Le = 1$ and $T_f = T_{ad}$. To further illustrate this effect, early measurements of the laminar burning velocities of spherical outwardly propagating flames, due to Palm-Leis and Strehlow,²⁷ are plotted as a function of flame radius in Fig. 2. These results are for propane-air flames at various fuel-equivalence ratios, ϕ . For these conditions, the laminar burning velocity increases with increasing flame radius when $\phi < 1.43$ and decreases with increasing flame radius otherwise. Since the experimental transition ϕ for T_f is near unity for propane-air flames, the fact that the experimental burning velocity changes its behavior for $\phi > 1$ supports the theoretical result that its transition Le also deviates from one.

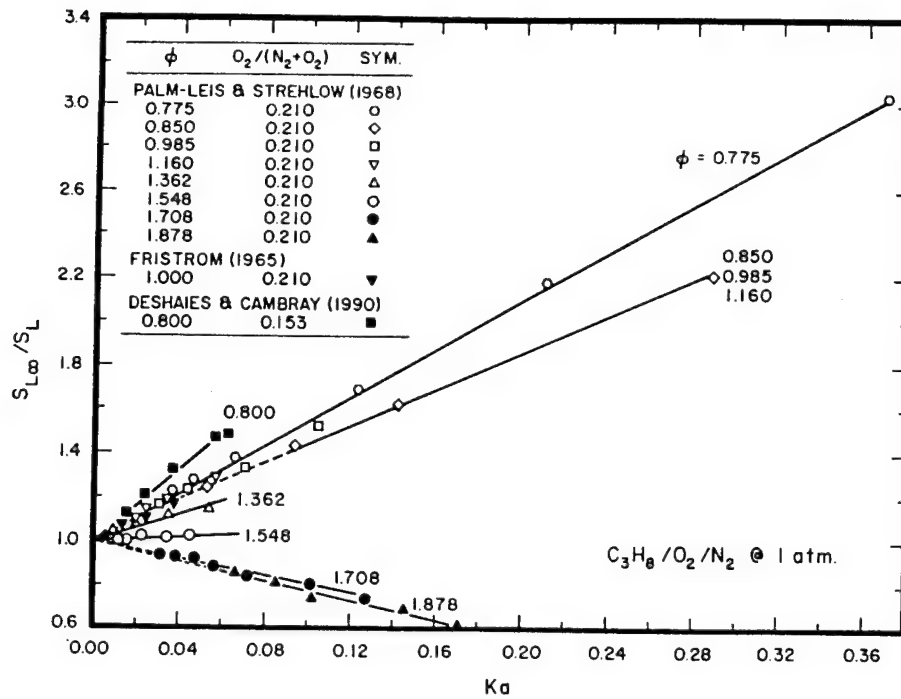


FIG. 3. Variation of laminar burning velocity with Karlovitz number for propane-air flames. Data from Palm-Leis and Strehlow,²⁷ Fristrom²⁹ and Deshaies and Cambray³⁰ as plotted by Kwon *et al.*²⁵

While theories^{15,20-23} of stretched flames yield basically the same result concerning the flame temperature response, differences exist concerning the predicted response for the laminar burning velocity. However, recent experimental measurements of the effects of flame stretch on laminar burning velocities using outwardly-propagating spherical flames,^{25,26} have been found to correlate conveniently, based on an early proposal of Markstein,²⁸ as follows:

$$S_L = S_{L\infty} - L_M K \quad (14)$$

where $S_{L\infty}$ is the laminar burning velocity for an unstretched flame ($K = 0$) and L_M is a measure of the response of the flame to stretch called the Markstein length. Then, basing the characteristic flame thickness on a characteristic mass diffusivity of the unburned gas, $\delta_L = D_u/S_L$, and introducing the Karlovitz number, Eq. (14) becomes:^{25,26}

$$S_{L\infty}/S_L = 1 + Mk Ka \quad (15)$$

where Mk is the Markstein number which is defined as the Markstein length normalized by the *current* characteristic flame thickness, that is, the flame thickness at the local stretched condition, as follows:

$$Mk = L_M/\delta_L = L_M S_L/D_u. \quad (16)$$

The Karlovitz number in Eq. (15) is defined in a corresponding manner, based on *current* flame properties, as follows:

$$Ka = KD_u/S_L^2. \quad (17)$$

The stretched burning velocities of propane-air mixtures are plotted according to Eq. (15) in Fig. 3,

from Kwon *et al.*²⁵ Results are shown for the measurements of Palm-Leis and Strehlow,²⁷ Fristrom,²⁹ and Deshaies and Cambray.³⁰ (Note that questions raised in Ref. 25 about the Palm-Leis and Strehlow²⁷ data have been resolved by recent measurements,²⁶ as discussed later.) It is seen that over the range of the measurements, which do not approach either quenching conditions with $Ka = 0(1)$,¹⁵ or flammability limits, the correlation between $S_{L\infty}/S_L$ and Ka is linear for a given ϕ , implying a constant Mk from Eq. (15). As pointed out earlier, the neutral-stability condition, $Mk = 0$, is reached near $\phi = 1.4$, with laminar burning velocities decreasing with increasing Ka ($Mk > 0$) for $\phi < 1.4$ and increasing with increasing Ka ($Mk < 0$) otherwise.

The large variations of laminar burning velocities with stretch is a striking feature of the results of Fig. 3, for example, $S_{L\infty}/S_L$ varies in the range 0.6–3.1 even though $Ka < 0.37$ for the test range. Applying Eq. (15), Mk ranges from -2.2 at $\phi = 1.878$ to 5.5 at $\phi = 0.775$ for the measurements of Palm-Leis and Strehlow;²⁷ notably, recent results for $H_2/O_2/N_2$ and hydrocarbon-air flames also yield comparable values of Mk .^{25,26} These large values of Mk show that effects of preferential diffusion cause laminar premixed flames to be very sensitive to modest levels of stretch, even well away from quenching conditions: this helps explain the importance of preferential-diffusion effects for strongly turbulent flames discussed earlier (see Fig. 1 and Refs 9–12). Additionally, Mk is significantly greater than unity for hydrocarbons at $\phi \leq 1$ (methane is an exception

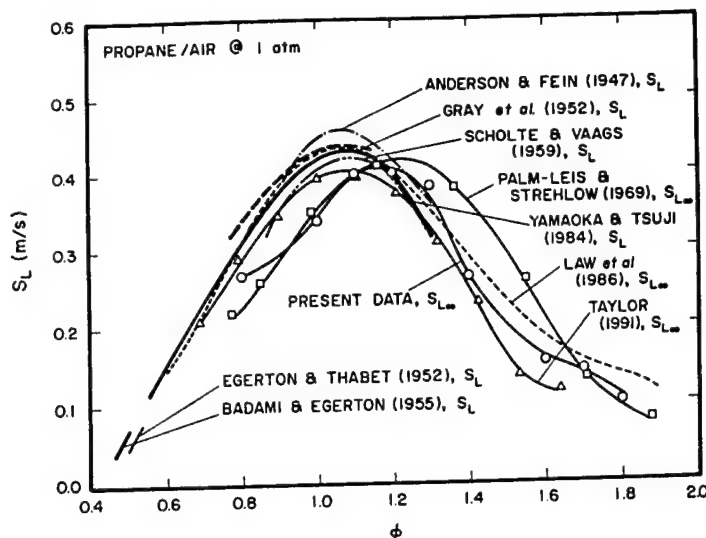


FIG. 4. Laminar burning velocities as a function of fuel-equivalence ratio for propane-air flames at normal temperature and pressure as plotted by Tseng *et al.*²⁶

where unstable conditions are in the range $\phi < 0.7^{26}$, suggesting strong effects of preferential-diffusion stability on flamefront properties for practical applications at these conditions, for example, furnaces, automobile engines and aircraft propulsion systems.

The large effects of stretch on laminar burning velocities also raise questions about existing data in the literature for this important fundamental property. In particular, the laminar burning velocity for an unstretched (planar) flame, $S_{L\infty}$, is the fundamental property of interest, however, in spite of efforts to minimize and correct for stretch, the database generally involves finite and unspecified values of Ka . Thus, effects of stretch combine with other experimental uncertainties to yield considerable uncertainties about $S_{L\infty}$.

An indication of potential problems with existing measurements of fundamental laminar burning velocities can be obtained by comparing measurements from various sources. Results of this type for methane, ethane, ethylene and propane-air flames are presented by Tseng *et al.*²⁶ a typical example for propane-air flames is illustrated in Fig. 4. Results that are plotted include measurements for outwardly propagating spherical flames from Tseng *et al.*,²⁶ Palm-Leis and Strehlow²⁷ and Taylor³¹ (the latter two re-analyzed in Ref. 26 as discussed in connection with Fig. 3), measurements for stagnation point flames from Law and coworkers³²⁻³⁵ and Yamaoka and Tsuji,³⁶ and several other measurements using a variety of techniques: original sources cited in Ref. 26 should be consulted for the details of these experiments. The outwardly-propagating spherical flame results of Tseng *et al.*,²⁶ Palm-Leis and Strehlow²⁷ and Taylor³¹ have been corrected as discussed in connection with Fig. 3 in order to provide estimates

of $S_{L\infty}$. Similarly, the stagnation point measurements of Law and coworkers³²⁻³⁵ involve techniques where effects of stretch are quantified and subtracted out so that $S_{L\infty}$ can be estimated (see Ref. 35 for a complete compilation of measurements based on this approach). The remaining measurements have not been corrected for stretch.

The results illustrated in Fig. 4 are not very encouraging with respect to current knowledge concerning fundamental laminar burning velocities, even considering results where corrections for stretch have been made. Among the corrected measurements, the results of Palm-Leis and Strehlow²⁷ probably involve an error in reported fuel-equivalence ratios, as discussed in Ref. 26, and will not be considered further. The remaining measurements only are in fair agreement for ϕ near 1.4, which corresponds to near-neutral conditions for propane-air flames where values of Ma , and thus effects of stretch, are small (see Fig. 3 and Ref. 26). At both fuel-lean and fuel-rich conditions where Ma is large for propane-air flames,²⁶ there are substantial differences between the various measurements which are probably mainly attributable to effects of stretch and the procedure through which the stretch effects are subtracted out (see Refs 26, 32-35 and 38, and references cited therein, for discussions of this methodology). Clearly, resolving the uncertainties of laminar burning velocities must have high priority because this fundamental property impacts the interpretation of a variety of flame phenomena. Additionally, these uncertainties compromise the development of methods to numerically simulate premixed laminar flames using detailed chemical-kinetic and transport models because they generally are calibrated using laminar burning velocity data (see the recent review by Warnatz,³⁹ and references cited therein).

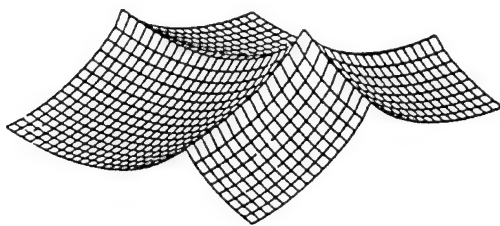


FIG. 5. Thermal-expansion-induced steady folds of fully-developed hydrodynamic instability. Predictions from Sivashinsky.⁴³

Measurements at μg offer opportunities to resolve problems of laminar burning velocities. For example, effects of buoyancy-induced stretch for the outwardly-propagating spherical flame technique can be eliminated in this manner, increasing the reliability of determinations of both $S_{L\infty}$ and Mk . Work along these lines has been carried out by Ronney and coworkers, and will be discussed in connection with near-limit phenomena, but much remains to be done. An alternative approach involves consideration of one-dimensional premixed flames around porous cylindrical and spherical burners at μg . This configuration is attractive because the flames are steady and the flow is normal to flame, which implies that the flames are stretchless.¹⁵ Work with cylindrical flames of this type has been initiated.⁴⁰

Other implications of effects of stretch on reaction intensity, flamefront instability and flammability limits are addressed in the following.

3.3. Flamefront Instability

Laminar premixed flames do not always exhibit smooth flame surfaces. Various types of instabilities can develop over flame surfaces, causing wrinkling and fundamental changes of flame topography and structure.^{25,26,41-48} When effects of instabilities are present, the burning intensity varies over the flame surface, leading to local regions of high reactivity (or hot spots) as well as extinguished regions through which reactants can leak. It has also been suggested that flamefront instabilities could lead to self-turbulization of flames.⁴³ Other interactions between stability phenomena and turbulent flamefronts have already been discussed in connection with Fig. 1.

As briefly noted earlier, there are three major mechanisms of flamefront instability of interest to laminar and turbulent premixed flames: hydrodynamic, preferential-diffusion and Rayleigh-Taylor instabilities. Hydrodynamic (or Landau-Darrieus) instability involves the growth of disturbances due to pressure fields caused by density changes across the flame surface. The linear stability analysis of Landau and Darrieus treats the flame like a structureless density discontinuity propagating through an other-

wise incompressible medium and shows that the flame is unstable to perturbations of all wavelengths.²⁸ This analysis presents a dilemma, however, because smooth flames are routinely observed in the laboratory. A nonlinear analysis has shown that such instabilities will evolve into folds having substantial dimensions relative to the disturbances as illustrated in Fig. 5.⁴³ This mechanism has been observed for large-diameter spherical laminar flames,^{25,44} but it is probably not of primary interest for either laminar premixed flame studies, where flame sizes are generally smaller, or most practical applications involving turbulent flames, where momentum fluctuations from turbulence tend to dominate the distortion of flame surfaces.

Preferential-diffusion instability is the mechanism of major interest for turbulent flames, due to its capability to enhance or retard turbulent distortion of the flame surface in the laminar flamelet regime.^{5-12,19} Two classes of preferential-diffusion instabilities have been analyzed in the past: diffusive-thermal and diffusive-diffusive instabilities. The diffusive-thermal instability mechanism was first described by Zel'dovich and coworkers,^{45,46} and has received substantial subsequent theoretical attention using activation-energy asymptotics of a one-reactant flame.^{18,43,47} Diffusive-thermal instabilities involve preferential diffusion of mass and heat, affecting flame temperatures as discussed in connection with Eq. (9), and accordingly the local burning velocities. In this case, the flame bulges into the reactant yield $Ka > 0$, similar to freely propagating spherical flames, while the flame bulges into the combustion products yield $Ka < 0$, similar to the tips of Bunsen burner flames.¹⁵ Then for $Le < 1$, bulges into the reactant (product) have $T_f/T_{ad} > 1$ (< 1) through Eq. (9), causing the bulges to grow and the flame to be unstable. The opposite behavior occurs for $Le > 1$. Thus, fuel-lean premixed flames of heavy hydrocarbons are stable, as noted earlier, because the fuel is the deficient reactant and its large molecular weight implies a low mass diffusivity so that $Le > 1$. More detailed analysis shows that diffusive-thermal effects are stabilizing for small wavelengths, thereby partially resolving the Landau-Darrieus dilemma.^{43,47} At longer wavelengths, the diffusive-thermal effect is destabilizing for small Le , leading to the formation of cellular flames, stabilizing for intermediate values of Le , and destabilizing for large Le , leading to pulsating flames or travelling waves over smooth flame fronts. A nonlinear analysis further shows that the cells evolve in time in a chaotic manner (see Fig. 6.⁴³).

Diffusive-diffusive instability was first described by Manton *et al.*:⁴⁸ it involves the preferential diffusion of reactant species. Figure 7 is a sketch illustrating the mechanism, involving a perturbed thin flame propagating at its local burning velocity with respect to the unreacted mixture. The flame surface is a sink for reactants; therefore, the relative mass flux of the

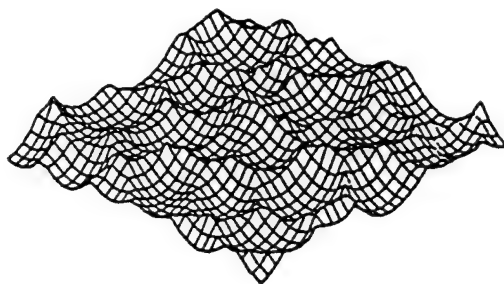


FIG. 6. Diffusive-thermal unstable cellular flames in a state of chaotic self-motion. Predictions from Sivashinsky.⁴³

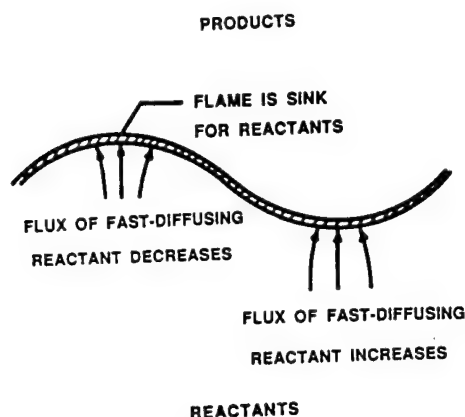


FIG. 7. Sketch of the diffusive-diffusive preferential-diffusion instability mechanism.

fast-diffusing reactant increases at bulges projecting into the reactant and decreases at bulges projecting into the combustion products, modifying the reactant mixture ratio at bulges accordingly. If the laminar burning velocity increases (decreases) with increasing concentrations of the fast-diffusing reactant, the bulges projecting into the reactant have larger (smaller) burning velocities than the bulges projecting into the product so that the bulges grow (decay) yielding an unstable (stable) flame. For example, hydrogen is the fast-diffusing reactant in $H_2/O_2/N_2$ flames which have a maximum burning velocity near $\phi = 1.8$; therefore, these flames should be unstable for $\phi < 1.8$ and stable for $\phi > 1.8$ according to diffusive-diffusive stability theory.¹⁰⁻¹² However, diffusive-diffusive theories do not predict the transition condition between stable-unstable conditions very well since recent observations find this transition near $\phi = 1.4$.²⁵

The deficiencies of simplified diffusive-thermal and diffusive-diffusive instability mechanisms arise because preferential-diffusion instability generally involves combined effects of preferential diffusion of the various species with respect to one another and with respect to thermal energy. This combined behavior can be related to effects of stretch through Eq. (15) if Mk is known. For $Ka > 0$, Mk can be found from results such as Fig. 3. In contrast, $Ka < 0$ is

problematical due to the difficulties of reliably measuring S_L for these conditions. On the other hand, limited existing measurements suggest that plots for $Ka > 0$ at a particular ϕ can be extrapolated at least qualitatively into the region where $Ka < 0$.³⁰ Thus, bulges into the reactant have $Ka > 0$ so that for $Mk < 0$ (> 0) the burning velocity increases (decreases), the bulges grow (decay), and the flame is unstable (stable). However, additional theoretical and experimental studies are needed to resolve preferential-diffusion behavior and the properties of Mk when $Ka < 0$ and when flammability limits and quenching conditions are approached. As discussed earlier, experiments at μg are particularly valuable to study behavior near limits and quenching conditions, where S_L is small in comparison with the criterion of Eq. (4); existing results of this type will be discussed in Section 3.5.

Rayleigh-Taylor instability involves accelerations normal to a density discontinuity like a flame.²⁸ In general, acceleration of the heavier fluid toward (away from) the lighter fluid is unstable (stable). This kind of instability has been observed for flames in shock tubes, where gravity has little effect,²⁸ but it is most significant for gravitational acceleration. Thus, upward (downward) propagating flames, where the heavier reactant gas is above (below) the lighter combustion products, are unstable (stable) to gravitationally-induced Rayleigh-Taylor instability. Such instabilities yield a convex shape to upward propagating flames and a somewhat planar shape for downward propagating flames, for flames that are otherwise stable.^{49,50} This mechanism influences burning velocities and limits, and obscures other instability mechanisms at ng . Eliminating the gravitationally-induced Rayleigh-Taylor instability is an additional motivation for combustion experiments at μg .

Studies have shown that the gravity-induced Rayleigh-Taylor instability interacts strongly with the preferential-diffusion instability. The classical example⁵¹ is shown in Fig. 8 which is an illustration of the observed flame shapes of upward propagating flames in very lean hydrogen-oxygen mixtures with inerts having different molecular diffusivities. It is seen that the flame can either form a closed surface covered by brightly luminous streaks (Fig. 8a), or is composed of a large number of flamelets rising in a zig-zag motion (Fig. 8b), or assumes a 'jelly-fish' like structure (Fig. 8c), as the mixture mass diffusivities are varied. The corresponding downward propagating flames do not exist in these situations because the mixtures are too lean.

Kailasanath and coworkers⁵²⁻⁵⁸ have completed several computational studies of the interactions between Rayleigh-Taylor and preferential-diffusion instabilities. These studies considered a flame tube configuration, like that of Strehlow *et al.*,⁵⁹ where flames propagate upward or downward at ng , or propagate at μg (taken to be 0 g). The computations

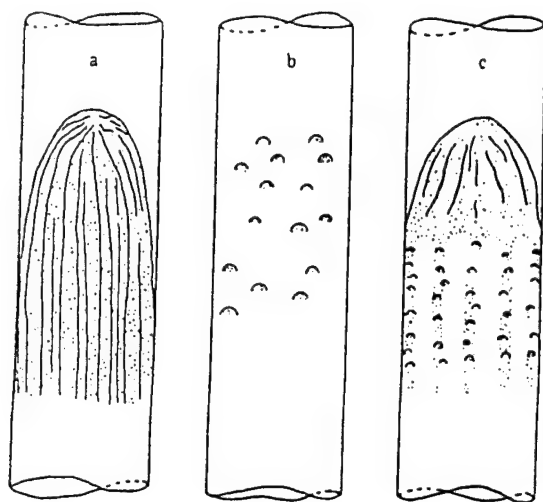


FIG. 8. Sketches of lean hydrogen flames propagating upward in a closed vertical tube, demonstrating the effects of preferential diffusion interacting with Rayleigh-Taylor instability. Reactant concentrations by volume are: (a) 6% H_2 , 94% dry air; (b) 6% H_2 , 20% O_2 , 74% SO_2 ; (c) 6% H_2 , 84% O_2 , 10% $SiCl_4$. From Böhm and Clusius.⁵¹

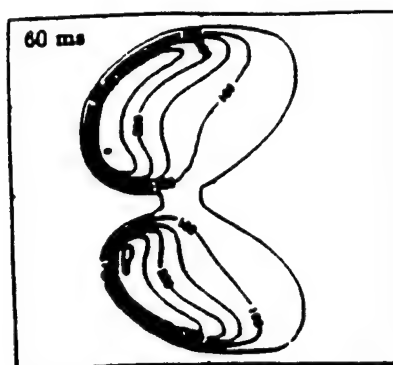
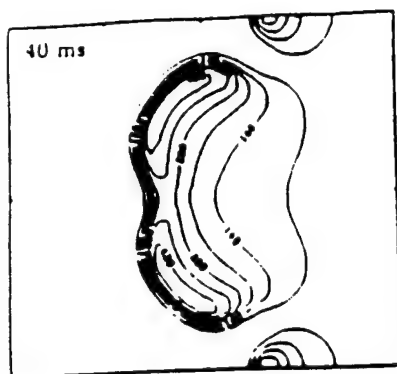
were limited to the two-dimensional, time-dependent propagation of flames in $H_2/O_2/N_2$ mixtures, with chemistry involving 24 reversible reactions, diffusion using Fick's law, and consideration of variable thermochemical and transport properties. The initial conditions were found by computing the properties of a purely one-dimensional flame and then perturbing the solution by displacing the central portion toward the reactant side of the flame.

Typical results from the computations of Kailas-anath and coworkers⁵²⁻⁵⁷ are illustrated in Figs 9 and 10. The premixed flame sheet is indicated by contours of the radical OH because radical species generally are confined to regions where chemical reaction is significant, for example, near the hot boundary of the flame surface. The contours are illustrated at various times after perturbation of the flame surface for upward and downward propagation at ng and at 0 g, for lean H_2/O_2 flames that are subject to preferential-diffusion instability. Conditions illustrated in Fig. 9 are for a rapidly propagating flame with strong preferential-diffusion instability. In this case, the effects of Rayleigh-Taylor instability are relatively unimportant during the period of the computations (up to 60 ms), although they would become important at longer times. The effects of preferential-diffusion instability are seen by the flame surface breaking up into cells with negligible reaction between the cells due to both reduced H_2 concentrations and temperature levels caused by stretch within the cusp-like regions projecting into the combustion products. Depending upon the mixture ratio, the time of propagation and the width of the computational domain, the cells can divide and their number will vary.

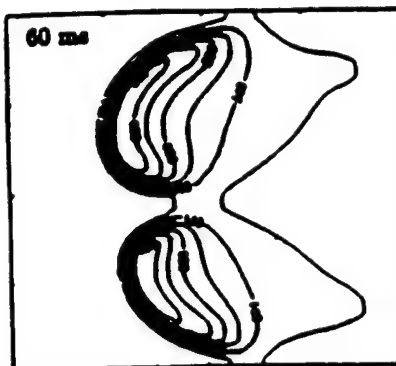
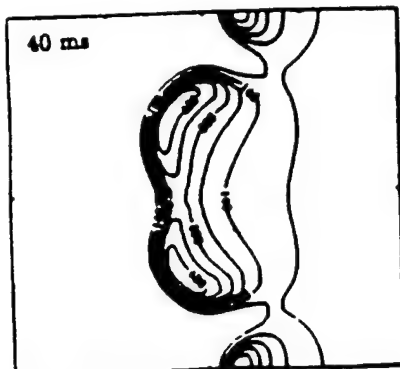
The effect of Rayleigh-Taylor instability at ng is much more dramatic for the more slowly propagating flame illustrated in Fig. 10. To begin with the 0 g results, the cell structure due to preferential-diffusion instability is qualitatively similar to Fig. 9; however, evolution of the cell structure is slower due to the lower reactivity of this flame. For upward propagation, however, Rayleigh-Taylor effects remain about as strong as before so that a strong buoyant cell is produced. Regions of negative stretch near the cusps into the burned gas cause the flame to quench or extinguish, behavior that also is associated with flame extinction due to buoyancy near the flammability limits at ng.¹⁵ Conversely, Rayleigh-Taylor effects act to stabilize the flame during downward propagation at ng, completely eliminating the growth of diffusional instabilities for the conditions of Fig. 10. More extended calculations for the conditions of Fig. 10 show that preferential-diffusion instability cells are overwhelmed by Rayleigh-Taylor effects at ng: upward propagating flames evolve into a bubble-like surface while downward propagating flames oscillate between mildly concave and convex flame surfaces.⁵³ These studies have also shown that the diffusive-thermal mechanism has a much stronger effect on flamefront instability than the diffusive-diffusive mechanism for this reactant mixture.

The work described above⁵²⁻⁵⁷ clearly establishes the importance of diffusive instabilities near limits. A recent study⁵⁸ has further shown that chemical oscillation can also occur for near-limit rich hydrogen-air flames, causing pulsating instability. Specifically, as the hydrogen concentration in a rich mixture is gradually increased for a planar flame, damped and then undamped oscillations in the flame velocity appear (Fig. 11). This persists until the mixture becomes too rich to support flame propagation. This oscillation is thought to be caused by the crucial $H-O_2$ branching-termination chain, although the precise mechanism has not been explained. The potential influence of such chemical instability on flamefront cellular instability, especially for near-limit mixtures, merits further study. Furthermore, the chemical-kinetic and transport aspects of the computations described above involve more approximations than contemporary simulations of this type,^{33,60} and real-world processes are three dimensional. Thus, more work is needed, involving both numerical predictions as computational capabilities increase and experiments to provide direct measurements of preferential-diffusive instability effects at μg . These results will be useful for developing and evaluating approximate numerical predictions of these processes. In particular, because buoyancy interactions are most prominent for near-limit mixtures, experiments at μg eliminate the buoyancy interaction and allow detailed study of the preferential-diffusion effects that are most relevant for practical premixed flames.^{1,42,61-63}

Upward



ZERO-G



DOWNWARD

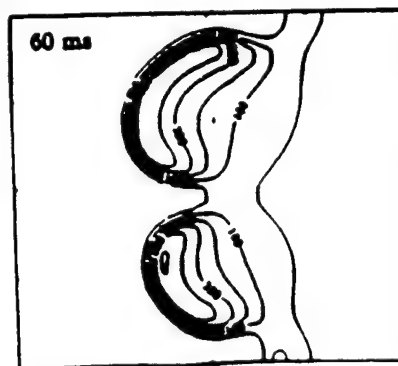
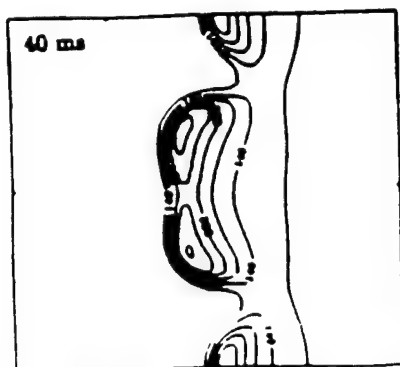


FIG. 9. OH radical number density contours ($\times 10^{14}$) for rapidly propagating H_2/O_2 flames ($H_2:O_2:N_2/1.5:1:10$ mixture by volume). Reactants are toward the left-hand side. Predictions from Kailasanath *et al.*⁵²

3.4. Flammability Limits

Flammability limits are a fundamentally and practically important concept in combustion.^{41,42} Empirically, it has been found that sufficient dilution of a combustible mixture with excess fuel, oxidant, or inert gas can render the mixture nonflammable. The critical composition at which this occurs is the flammability limit. When there is a fixed relationship between the concentrations of the inert gas and the fuel or oxidant, for example, the fixed relative oxygen

and nitrogen concentrations in air, there are only two flammability limits at a particular pressure and temperature: the lean flammability limit where there is excess oxidant and the rich flammability limit where there is excess fuel. Naturally, a knowledge of the lean and rich flammability limits of a given fuel in air is important for assessing fire and explosion hazards in, for example, mine galleries, chemical refineries, and manned spacecraft. This information also is relevant to the design of lean-burn engines and the use of low-BTU gases.

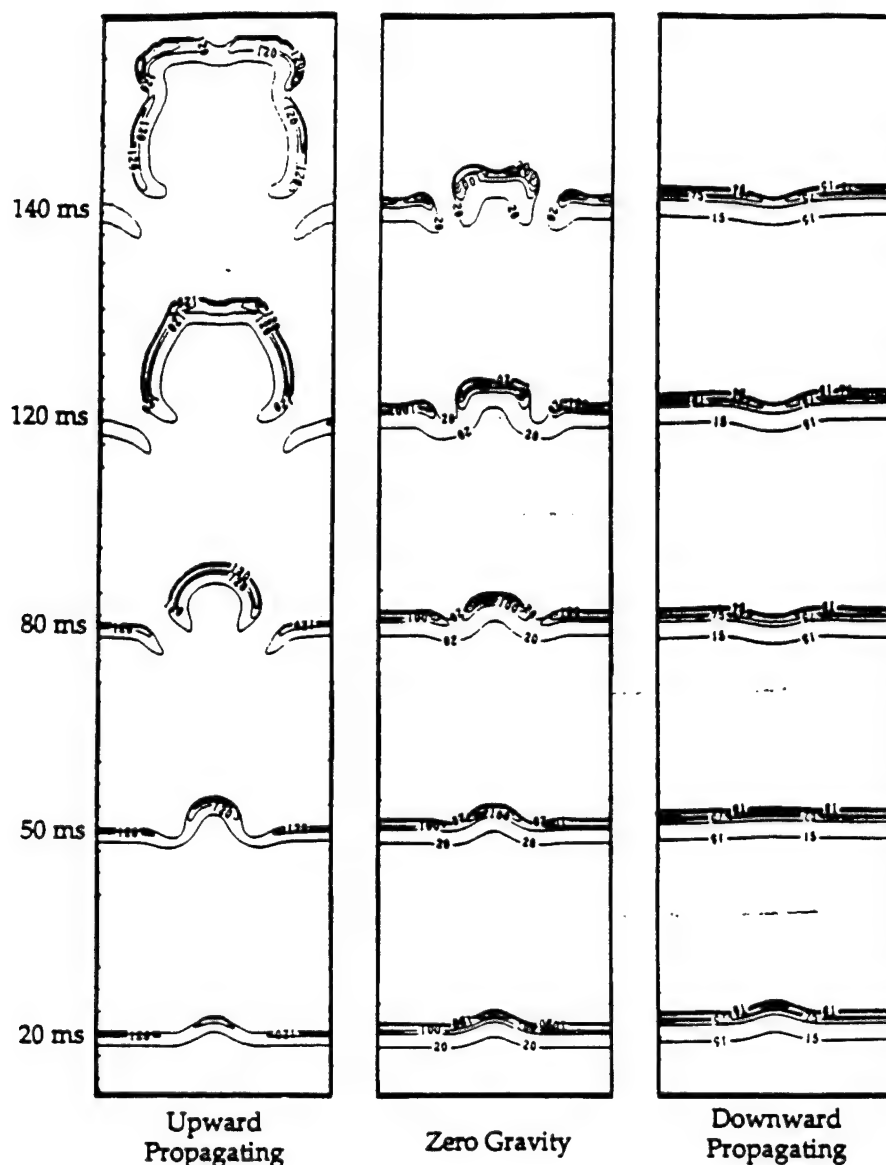


FIG. 10. OH radical number density contours ($\times 10^{14}$) for near-limit H_2/O_2 flames (10% H_2 /air by volume). Reactants are toward the top. Predictions from Patnaik *et al.*⁵³

Studies of flammability limits generally are concerned with the numerical values of these limits and the mechanism(s) responsible for their existence. Flammability limits determined at ng exhibit significant effects of buoyancy, causing much wider limits for upward-propagating flames than for downward-propagating flames, and limits are different from both at μg . Experimental studies addressing this issue include Strehlow *et al.*⁵⁹ and Ronney and coworkers.⁶⁴⁻⁷² Strehlow *et al.*⁵⁹ used a configuration where the flame is ignited by a spark and propagates inside a tube, either upward or downward at ng or μg , similar to the results illustrated in Figs 9 and 10. Ronney and coworkers⁶⁴⁻⁷⁰ consider spark-ignited flames propagating from the center of a chamber, in an initially still gas, as a more or less spherical flame.

In the latter work, upward and downward propagation limits at ng were defined differently than for tubes: the upward limit implied flame propagation to the top of the vessel, the downward limit implied flame propagation throughout the vessel. These measurements employed a drop tower to reach μg , except for recent experiments in aircraft.⁶⁹

Typical lean flammability limits at ng and μg , from Ronney and Wachman,⁶⁴ are plotted as a function of pressure in Fig. 12. The results are for lean methane-air flames so that $\text{Mk} < 0.26$. The results indicate that flammability limits at μg generally fall between those for downward and upward propagation at ng: downward propagation is the most difficult because buoyancy sweeps the flame to the top of the chamber at the low burning velocities near

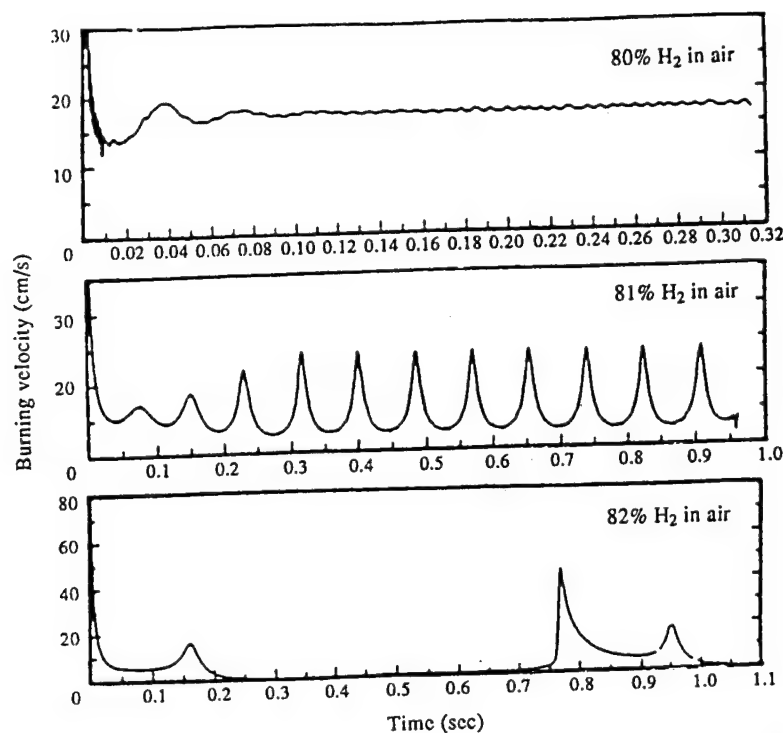


FIG. 11. Laminar burning velocity as a function of time, illustrating oscillatory near-limit behavior for rich hydrogen-air flames. From Kailasanath *et al.*⁵⁸

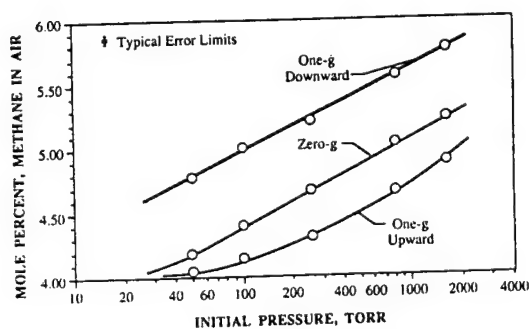


FIG. 12. Lean flammability limits of methane-air mixtures as a function of pressure for upward and downward propagation at ng and μg . From Ronney and Wachman.⁶⁴

limits so that the chamber walls quench the flame. The different flow conditions and flammability definitions for the flame tube experiments of Strehlow *et al.*⁵⁹ yielded somewhat different results: lean limits for methane-air mixtures at atmospheric pressure were 5.25, 5.25 and 5.85% for upward, μg and downward propagation, which are higher with less difference between upward and μg propagation than for the results shown in Fig. 12. Additionally, Strehlow *et al.*⁵⁹ find conditions where lean limits at μg are outside those at ng; for example, propane-air flames at atmospheric pressure had lean limits of 2.15, 2.06 and 2.20% for upward, μg and downward propagation. This probably occurs because $Mk > 0$ for lean propane-air flames, as seen in Fig. 3, so that posi-

tively stretched upward propagation of the propane-air flames causes reduced burning velocities and flame temperatures, in contrast with the lean methane-air flames where $Mk < 0$ and burning velocities and flame temperatures increase.²⁶ Then increased stretch rates for upward propagation at ng in comparison with μg , due to buoyant motion, causes a correspondingly higher lean flammability limit. The same argument implies that rich flammability limits for methane-air mixtures, where $Mk > 0$, would be broader at μg than at ng, but measurements in this region have not been reported.

Other observations supporting effects of stretch on near-limit behavior will be discussed subsequently. However, the broader lean flammability limits at μg for reactants with $Mk > 0$ have important implications for spacecraft fire and explosion hazards because most lean hydrocarbon-air flames are in this regime, as noted earlier. Thus, any notion that flammability limits for gases at ng are applicable to conditions at μg is erroneous. Additional measurements of lean flammability limits for a wider range of reactants are clearly needed to provide the technology base required for safe spacecraft operation with respect to fires and explosions.

The fact that flammability limits do seem to exist at μg resolves a controversy regarding the role of buoyancy in limit phenomena. That is, previous theories relating such limits to natural convection predict that no limits should exist in the absence of gravity. Thus, the μg experiments that have just been dis-

cussed have ruled out buoyancy as a controlling mechanism for the occurrence of flammability limits.

There also have been disagreements about whether the flame should have a finite or vanishing speed at the flammability limit. Previous studies cannot answer this question satisfactorily due to the large buoyant flow velocities or heat loss to the burner. Recent μg experiments, however, show that these flame speeds assume finite, albeit very small, values of around a few centimeters per second.^{66,67}

The fundamental controlling mechanism(s) of flammability limits are as yet unidentified, although valuable insights have been gained in some recent studies.^{23,73,74} Clearly, if the flammability limit is a viable concept, and its value is a unique property of a given combustible, then the mechanism(s) should be independent of external influences and thereby predictable from first principles. At present it appears that there are two major classes of fundamental flammability limits that can be defined from first principles, depending on whether the flame is stretched or not. For the unstretched flame, the relevant phenomenon is the failure of propagation of the one-dimensional, planar flame in the doubly-infinite domain. The intrinsic, omnipresent mechanisms which can cause extinction of such a flame are radiative heat loss and chemical-kinetic chain termination. Radiative heat loss decreases the flame temperature which in turn exponentially reduces the heat generation rate in the flame. Extinction can be expected when the heat loss rate becomes excessive. On the other hand, chemical-kinetic flammability is based on the consideration that the two-body chain-branching reactions are temperature sensitive while the three-body chain-termination reactions are usually temperature insensitive. Thus, by continuously reducing the concentration of the lean reactant, the corresponding reduction of the flame temperature progressively weakens the intensity of the branching reactions relative to that of the termination reactions. It is therefore reasonable to expect that at a certain concentration the overall reaction rate would have been so weakened that the flame is readily extinguished by unavoidable perturbations in the system.

Recently, the separate concepts of flammability due to heat loss and chain termination have been unified through a numerical simulation of the non-buoyant (planar) flame in the doubly-infinite domain, with radiative heat loss and detailed chemistry, for methane-air and hydrogen-air flames.⁷⁴ It is shown that as the flammability limit is approached, the flame response exhibits an extinction turning point which is characteristic of extinction due to heat loss. Furthermore, near this turning point, the normalized sensitivity of the rate of the dominant chain-termination reaction to that of the dominant chain-branching reaction becomes $O(1)$, indicating that the chain-termination reaction starts to have a controlling influence on the flame response. These results therefore suggest that at the flammability limit,

the branching reaction is so weakened relative to the termination reaction that the overall heat release rate is rapidly reduced. Extinction occurs when the heat release rate cannot keep up with the radiative heat loss rate. The theory also shows that the laminar flame speed has a finite value at the flammability limit.

Since practically all realistic flames are subjected to stretch, it is then necessary to consider whether fundamental flammability limits can also be defined for stretched flames. For these stretch-affected limits the intrinsic mechanisms of heat loss and chain termination identified for the nonstretched flames should obviously still be operative. In addition, it is reasonable to conjecture that a unique stretch rate may also exist at such a limit. The flammability condition could therefore be expressed as a critical flame diameter for spherically propagating or stationary flames, or a critical strain rate for aerodynamically stretched flames. The possibility of defining flammability limits through spherical flames will be discussed next.

3.5. Near-Limit Phenomena

Due to their ability to sustain extremely weak flames having small propagation rates, μg experiments have identified some premixed flame phenomena that have not been previously observed. These include self-extinguishing flames (SEF) and stationary spherical flames (SSF). Both SEF and SSF are associated with conditions near flammability limits and involve preferential-diffusion effects. Thus, SEF and SSF have the potential to provide insight concerning the mechanism of flammability limits and to highlight the unique combustion properties of non-buoyant μg conditions.

3.5.1. Self-extinguishing flames

Ronney⁶⁵⁻⁶⁷ made the first observations of SEF during his experiments on outwardly propagating spherical flames near flammability limits. The nature of SEF can be seen from the plots of flame radius as a function of time for lean ammonia-air mixtures illustrated in Fig. 13. The two cases shown are for conditions just above and just below the lean flammability limit, both ignited using sparks having similar energies. This reactant system involves $Mk < 0$ with a progressively decreasing positive stretch as the flame radius increases. Thus, in agreement with the discussion associated with Eq. (9), and the results illustrated in Figs 2 and 3, the slope of the flame radius as a function of time, which is proportional to the burning velocity through Eq. (12), decreases with increasing radius as the flame temperature, and thus its reactivity, decreases. Propagation continues for the normal flame, with the burning velocity eventually approaching a constant value at large radii where the effects of stretch become small because the flame

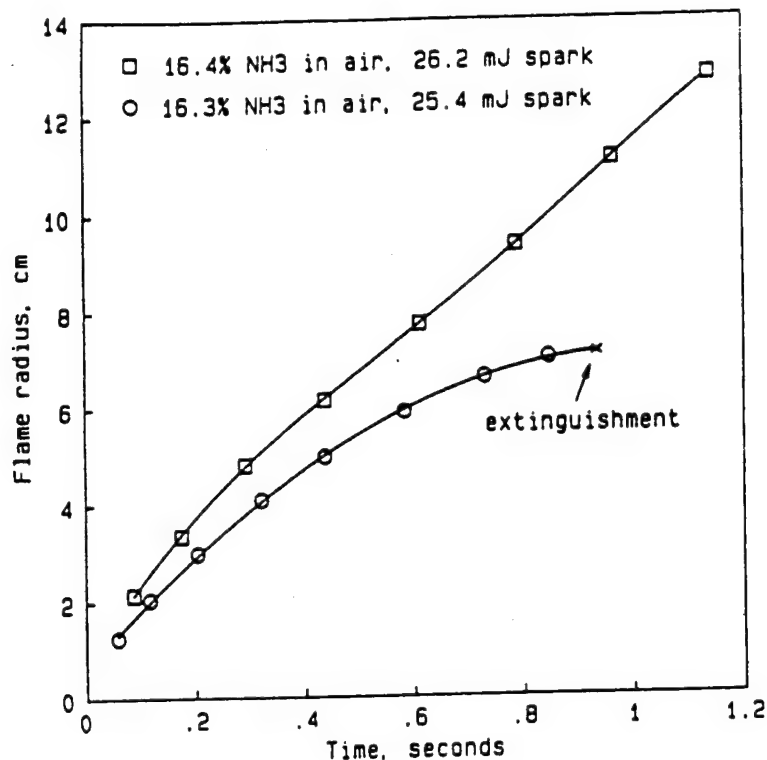


FIG. 13. Measured flame radius as a function of time for near-limit ammonia-air flames at μg . From Ronney.⁶⁷

is nearly planar. In contrast, propagation ends abruptly before a constant burning velocity is reached for the SEF. Notably, SEF are not caused by the added energy of the spark. Conventional nonpropagating flames normally consume some reactant due to the thermal disturbance of the spark before extinguishing, typically releasing reaction energies roughly 10 times larger than the spark energy. SEF, however, release 100–1000 times more chemical energy than the spark energy before extinguishing, so that their presence is not solely associated with initial overheating of combustion products by the spark.

Theoretical studies of SEF indicate that they are due to the interaction of stretch and radiative heat losses from the flame.^{67,70–72} First, SEF are observed for conditions where $Mk < 0$, for example, fuel-lean ammonia, methane or hydrogen–air flames, or fuel-rich propane–air flames. This suggests that the tendency for positive flame stretch to increase the flame reactivity for $Mk < 0$ compensates for the tendency of radiative heat losses to reduce flame temperatures. Since the effect of stretch decreases with increasing flame radius, radiative heat losses eventually dominate the process and the flame is quenched. Thus, for these conditions, the normal flammability limit can be attributed to thermal effects. However, whether such limits are a fundamental property of the reactants, which would be helpful for establishing universal limits, is still an open issue

due to the numerous effects that influence radiative heat losses from flames, for example, the dimensions of the combustion products, the radiative boundary conditions, etc.

Predictions of flame propagation and SEF behavior are currently based on approximate theories invoking simplified descriptions of the chemical and transport processes.^{67,70–72} Thus, their quantitative accuracy is limited. Nevertheless, the theories do provide support for the thermal mechanism of extinction, and represent the qualitative behavior of SEF. Typical predictions for lean methane–air flames are illustrated in Fig. 14 where the flame radius is plotted as a function of time for various mixtures near the flammability limit. By definition, all SEF operate outside the normal flammability limits. Additionally, reduced concentrations of the fuel near this lean limit (reduced ϕ) cause the SEF to extinguish at a progressively smaller radius. Thus, the normal lean limit is the flame where radiative heat losses just remain above critical levels over the entire region in which stretch is finite and flame reactivity exceeds the planar-flame limit.

Other features of SEF are also consistent with interactions between stretch and radiation. In particular, SEF are not observed when $Mk > 0$, which corresponds to $Le > 1$ within the context of the simplified theories of premixed flames, where the discussion of Eq. (9), or the results illustrated in Figs 2 and 3, indicate that flame temperatures, and thus

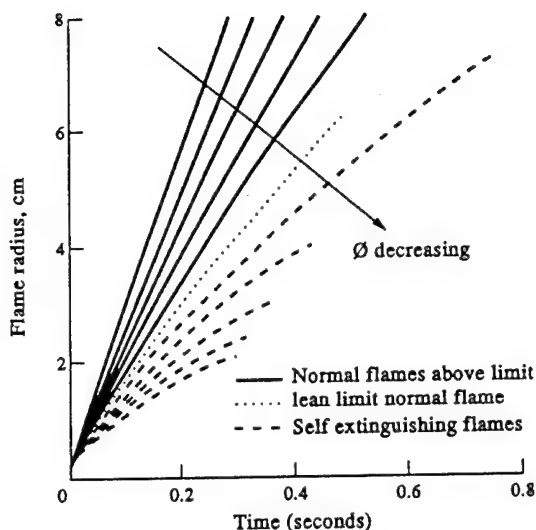


FIG. 14. Predicted flame radius as a function of time for near-limit normal flames and SEF in lean methane-air mixtures at μg . Compositions range from 5.6–4.6% methane in 0.1% steps ($\phi = 0.565$ –0.459). From Ronney.⁶⁶

burning velocities, increase with increasing flame radius. Then the flammability limit becomes strongly influenced by ignition conditions because once the flame is established the flame zone reactivity tends to increase until the flame surface becomes planar.⁶⁶ Due to the variation of ignition properties with the magnitudes and rates of spark energy deposition, whether flammability limits for these circumstances are a fundamental property of the reactants is also an open issue.^{41,42} The absence of intrusive effects of buoyancy at μg provides a useful environment to address this issue and should continue to receive attention by both experiments and analyses.

The previous discussion indicates that the study of SEF provides a new perspective on the interactions between preferential diffusion, stretch and radiative heat losses relating to flame propagation and limits. Notably, SEF behavior was never established at ng because buoyant motion strongly distorts these flames due to their low burning velocities (ca. 1–10 cm/s). Thus, experimental studies at μg clearly provide valuable new directions and insights concerning the properties of homogeneous premixed flames.

3.5.2. Stationary spherical flames

Another unique flame configuration experimentally observed at μg —the SSF—also provides valuable insights concerning premixed flame properties. SSF, often called flame balls, are observed in reactant mixtures near limits having very small Lewis numbers, such as lean hydrogen–oxygen flames. For such systems, preferential-diffusion instability occurs even near flammability limits causing the flames to break up into cells, similar to those illustrated in Figs 9 and 10. Examples from experiments at ng are illus-

trated in Fig. 8; this involves cap-shaped flamelets that rise through the mixture but do not seem to grow before being quenched by contact with the surfaces of test apparatus—behavior that was attributed to stabilization by convection.⁴¹ Similar flame systems are encountered at μg , however, where they develop into SSF whose radius remains constant, at least for the test times available for study at μg (up to 10 s during recent work).^{69,70} As will be discussed subsequently, the failure of SSF to continue outward propagation, and instead reach a stable fixed size, occurs due to heat losses in the burned gas. The mechanism involves radiation from the burned gas to the surroundings with steady temperatures maintained in the burned gas due to conduction (and perhaps some contribution from radiation) from the region of the flame surface. In a sense, this behavior is analogous to the stabilization of a premixed flame by heat losses near the surface of a flat flame burner, with the main differences between SSF and burner flames being the direction of energy flow from the flame to the stabilizing heat sink and the fact that SSF attain the unusual limit where mean velocities approaching the flame are zero.

The existence of SSF was proposed many years ago by Zel'dovich,⁴⁵ and subsequently studied by Buckmaster and Weeratunga,⁷⁵ however, SSF were never observed due to buoyant disturbances at ng . A sketch of the structure of these flames, based on current theories,⁷⁶ appears in Fig. 15. The conditions pictured involve a thin reaction zone at a radius, r_f , for an overall reaction that is complete, that is, all reactants become products. The mass fractions of reactants and products, Y_R and Y_P , the temperature, T , and the reaction rate per unit volume, \dot{w} , are plotted as a function of radius from the flame center, r . The reactant diffuses toward the flame, is converted to product within a thin reaction zone, and finally the product diffuses radially outward into the ambient environment. The SSF postulated by Zel'dovich⁴⁵ did not involve radiative heat losses. Recent analysis shows, however, that adiabatic SSF are unstable at their equilibrium radius, either collapsing or propagating outward for small disturbances.⁷⁶ Thus, radiative heat losses are depicted in Fig. 15, where the temperature decreases within the reacted core of the flame due to the balance between

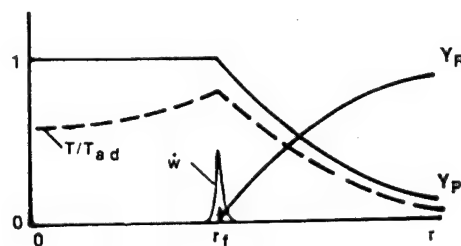


FIG. 15. Sketch of the structure of stationary spherical flames (SSF).

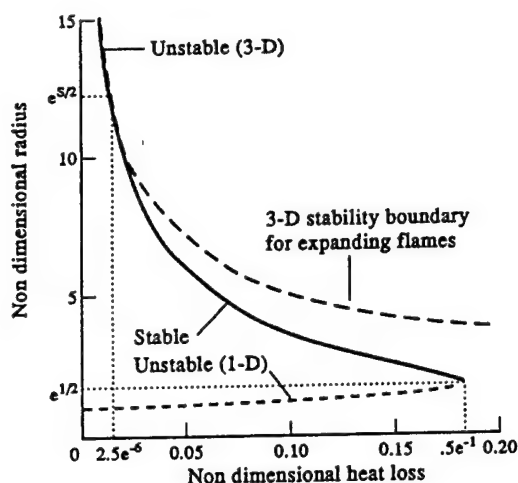


FIG. 16. Stationary dimensionless radius of SSF as a function of nondimensional heat loss. From Buckmaster *et al.*⁷⁶

radiation to the surroundings and heat conduction from the flame zone. Thus, the thermal energy release at the flame surface divides into two parts: one part flowing into the core to compensate for radiative heat losses to the surroundings, and the other part flowing directly outward toward the unburned gas similar to a conventional premixed flame.

The unique behavior of SSF has prompted analysis of conditions for their existence and stability.⁷⁵⁻⁷⁹ The initial theories,⁷⁵⁻⁷⁸ while simple in terms of chemistry and transport properties, yield predictions which are consistent with experimental observations. The predicted stability properties of the flames are illustrated in Fig. 16, in which the flame radius normalized by the radius of a corresponding adiabatic SSF, called the Zel'dovich radius, is plotted as a function of dimensionless heat loss (see Ref. 76 for definitions of these variables). There are two branches of the solution of flame radius as a function of heat loss. One is for a dimensionless flame radius less than $e^{1/2}$ which is unstable to one-dimensional (radial) disturbances. This branch includes the Zel'dovich SSF, which has a dimensionless radius of unity, highlighting the requirement for finite radiative heat losses to stabilize SSF. Similarly, the flames are unstable to three-dimensional disturbances for values of dimensionless radius greater than $e^{5/2}$. However, the branch of the solution with radii between $e^{1/2}$ and $e^{5/2}$ yields stable SSF with finite radiative heat losses similar to experimentally observed SSF.

Recently, a numerical simulation of SSF in hydrogen-air mixtures with detailed transport, chemistry and radiative loss has been conducted.⁷⁹ The simulation shows lean and rich limits beyond which no solution exists. It has been suggested that these limits can be identified as the corresponding flammability limits of the mixture. However, stability analysis of these numerically-generated flames, especially

the rich flame which is believed to be unstable based on Lewis number considerations from simplified analysis, needs to be conducted.

Due to the mathematical simplicity associated with the analysis of SSF, much can be learned about the structure and limits of flames by using SSF as a model flame problem. Furthermore, it has also been suggested that the appearance of SSF in limit situations could precede the onset of flaming combustion of fires in space environments. Thus, the study of SSF commands both fundamental and practical interest in μg combustion research.

4. GASEOUS NONPREMIXED FLAMES

4.1. Introduction

Studies of gaseous nonpremixed flames at μg have also largely been limited to laminar flames, even though most practical nonpremixed flames are turbulent. This is motivated by a desire to control the complexity of already complicated combustion processes, for experimental circumstances where diagnostics must be rudimentary by ng standards. However, the tactic also is justified to some extent by the evolution of laminar flamelet concepts for nonpremixed flames, somewhat analogous to those developed for premixed flames.

Ideas about laminar flamelet concepts were proposed for turbulent nonpremixed flames by Hawthorne *et al.*⁸⁰ not very long after the structure of laminar diffusion flames was described by Burke and Schumann.⁸¹ The main premise was that relationships between scalar properties in turbulent flames were the same as in laminar flames, that is, a turbulent flame corresponded to a distorted or wrinkled laminar flame. It also was assumed that preferential-diffusion effects were not significant, so that the scalar properties in laminar flames could be described by a variety of conserved scalars, related to the extent of mixing at a point. A popular choice for the conserved scalar is the mixture fraction, which is the mass fraction of elemental fuel species in a sample, independent of the current chemical composition. As a result, relationships between various scalar properties and the mixture fraction, called state relationships, can be found.⁸²⁻⁹¹ Then, predictions of the properties of mixture fractions in turbulent flames using a turbulence model, a large-eddy simulation, for example, yield all scalar properties—an approach called the conserved-scalar formalism.⁸²

The reason that state relationships can be found is that nonpremixed flames are often diffusion controlled with thin flame sheets, where the fuel and oxidant react to form combustion products. For such conditions, reaction rates are fast enough to approach local thermodynamic equilibrium at various local mixture fractions over a range of stretch

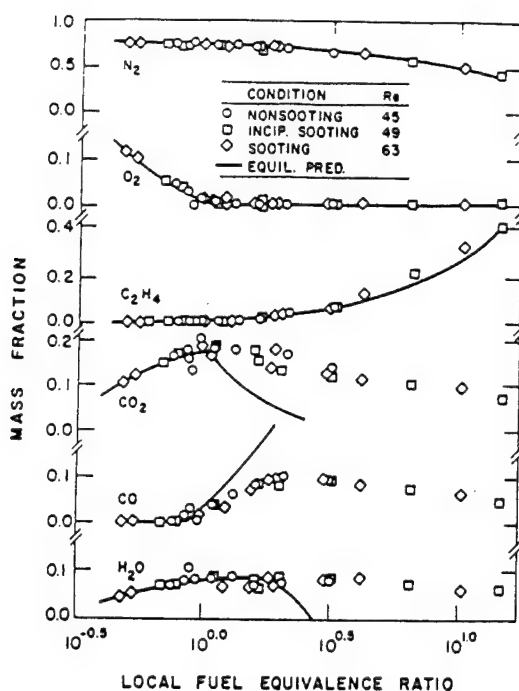


FIG. 17. State relationships for laminar ethylene-air flames. From Gore and Faeth.⁸⁴

rates. Thus, reactants involving simple molecules that have fast chemistry, for example, hydrogen-air and wet carbon monoxide-air flames, exhibit reasonably good state relationships, in spite of potential effects of preferential diffusion, that approach estimates assuming local chemical equilibrium for conditions not too near points of flame attachment or extinction.⁹⁰

The existence of state relationships for complex fuels such as hydrocarbons is more questionable due to potential fuel decomposition in the high-temperature region on the fuel-rich side of the flame sheet. However, Bilger⁸² noticed that major gas species in laminar hydrocarbon-air flames also satisfied state relationships. In this case, slow processes of fuel decomposition and soot chemistry caused departures from local thermodynamic equilibrium predictions for fuel-rich conditions; however, these departures are relatively universal over wide ranges of stretch rates. This behavior is illustrated in Fig. 17, where state relationships based on measurements in buoyant coflowing ethylene-air jet flames are shown along with predicted properties based on the assumption of local thermodynamic equilibrium.⁸⁴ These results are for various positions in the flames and burner exit Reynolds numbers, and thus a range of stretch rates, and still yield reasonably universal state relationships for these soot-containing flames. Subsequently, generalized state relationships have been developed for alkane and alkene-air flames, covering molar H/C ratios in the range 1–4, that exhibit relatively small effects of the wide variations of soot concentrations

within the flames.⁸⁶ Additionally, the use of state relationships has proved to be effective for estimating the structure and radiation properties of both nonluminous and luminous nonpremixed turbulent flames having modest stretch rates.⁸⁷

Similar to the laminar premixed flames, preferential diffusion could also modify the flame response through its influence on the flame temperature as well as through coupling with stretch.^{88,89} Such influences have not been addressed to the same extent as for premixed flames, possibly due to the fact that the influences are likely to be modest for diffusion flames because, in the limit of near-equilibrium combustion, the bulk flame response is basically decoupled from the reaction rate, which is sufficiently fast in any event, and hence only weakly dependent on the flame temperature. Furthermore, it has also been suggested⁹⁰ that preferential-diffusion effects are frequently being masked by experimental uncertainties even for flames involving H_2 . Thus, although the limitations of state relationships and laminar flamelet concepts are still being debated, and the fact that the methodology relies on the assumption of equidiffusion and also cannot treat finite-rate processes leading to pollutant emissions,⁹¹ the relevance of an understanding of laminar nonpremixed flames to turbulent nonpremixed flames of practical importance is widely accepted.

Studies of nonpremixed flames at μg have been limited to the laminar round jet flame. Although spherically symmetric diffusion flames would be simpler geometrically, and a burner-generated steady-state spherical flame is currently being planned for μg experimentation,⁴⁰ research with this configuration has centered on heterogeneous drop combustion which will be discussed later. Nevertheless, the jet flame is a fundamental configuration at μg due to the absence of complications from buoyancy and capabilities to vary stretch and reaction rates by varying burner diameters, jet exit velocities, fuel and ambient compositions, and the ambient pressure. Notably, simplified analyses of forced laminar jet diffusion flames have been available for some time,^{81,92} although measurements needed to evaluate predictions have only appeared recently.^{93–109} Results concerning the structure, stability and soot properties of these flames will be considered in the following sections.

4.2. Flame Structure

The earliest measurements of jet diffusion flames emphasized effects of gravity on their structure and stability properties in terms of extinction and stability.^{93–99} These experiments involved round fuel jets directed vertically upward in still air, with the flame ignited by a spark. In some instances, provision was made for a coflowing air stream around the fuel jet.⁹⁹ Tests were conducted at ng and μg , the latter using drop-tower facilities.

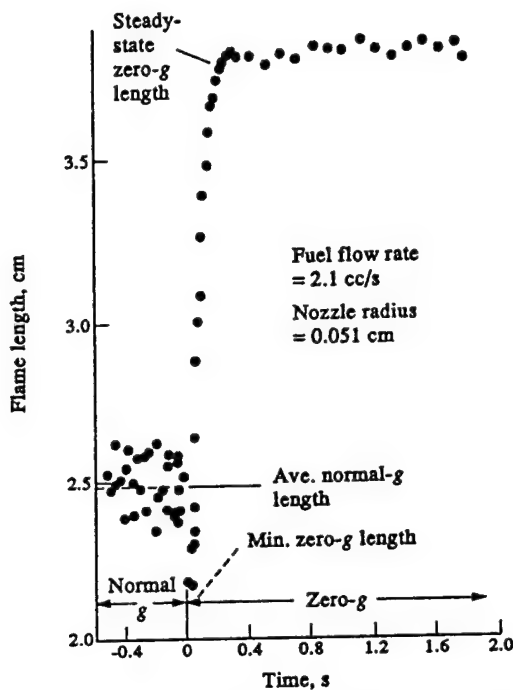


FIG. 18. Length of a methane-air laminar jet diffusion flame as a function of time at μg (near-steady conditions). From Haggard and Cochran.⁹⁶

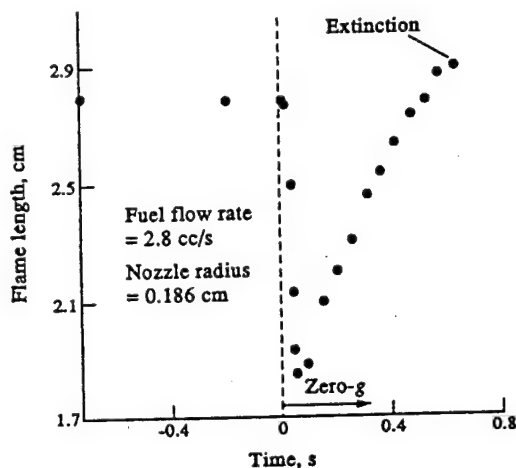


FIG. 19. Length of a methane-air laminar jet diffusion flame as a function of time at μg (extinguished conditions). From Cochran and Masica.⁹³

Some typical measurements of the properties of jet flames at μg are illustrated in Figs 18 and 19. These tests involved igniting the flames at ng and then releasing the apparatus to reach μg . The adjustment of the flame to μg causes it to become much shorter for a time and then to gradually lengthen toward its nonbuoyant configuration. In some instances, the flame extinguishes during this transient development process (Fig. 19). This behavior has been attributed to the accumulation of combustion products in the

flame region, due to loss of the buoyant convective flow, which excessively limits diffusion of oxygen to the flame sheet—particularly at the points of flame attachment near the jet exit.^{101,103} This view is supported by observations that even small levels of air coflow serve to sustain flames that would normally extinguish in the absence of coflow during transition to μg .⁹⁹ For conditions where the flame does not extinguish after entering μg , its length increases and eventually approaches a constant value that is consistent with steady flame structure (Fig. 18).

Although theories of nonbuoyant jet diffusion flames have been available since the original studies of Burke and Schumann,⁸¹ evaluation of predictions even for this fundamental flame configuration has been hampered by the complications of buoyancy at ng . The increasing availability of flame structure measurements at μg is beginning to rectify this situation, however, allowing theories of varying complexity to be evaluated.^{95,100,108-110}

The earliest models of nonbuoyant jet diffusion flames, aside from simple constant-property treatments of the qualitative features of the flow,⁹² are typified by the models of Edelman *et al.*⁹⁵ and Klajn and Oppenheim.¹⁰⁰ Major assumptions of these models are functionally the same: boundary-layer flow, neglect of radiation, thin flame sheet, equal diffusivities of all species and heat, and state relationships developed assuming complete oxidation of fuel species to products at the flame sheet. Other details differ somewhat, particularly the state relationships (see the original sources for more complete information). Bahadori *et al.*¹⁰⁴ report an evaluation of the models of Refs 95 and 100, based on flame length measurements at μg for methane, propane, ethylene and propylene flames burning in air. The results are illustrated in Fig. 20 where flame lengths are plotted as a function of the burner exit Reynolds number. The range of burner exit Reynolds numbers for these results is sufficiently high so that flame lengths increase nearly linearly with Reynolds number, in accord with simple boundary layer treatments of jet diffusion flames.⁹² Additionally, flame lengths in this regime are largely controlled by stoichiometric fuel-oxidant mass ratios, which are not very different for the fuels considered in Fig. 20. The two predictions are similar to each other and yield good results except for propylene-air flames that are considerably shorter than predicted. This difficulty is attributed to larger effects of radiative heat losses from the propylene flames.¹⁰⁴ This is plausible because propylene-air flames are known to have considerably higher radiative heat losses than the other fuels tested (see Ref. 87 and references cited therein), and increased radiative heat losses tend to decrease flame lengths.^{108,109} However, the effect seen in Fig. 20 is surprisingly large, and experimental difficulties may also be a factor.

More recent theories have addressed the limita-

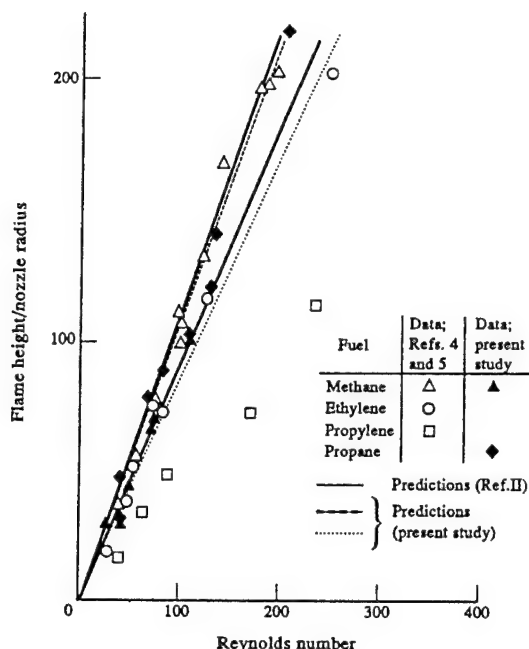


FIG. 20. Predicted and measured flame lengths of nonbuoyant laminar jet flames. Measurements from Haggard,⁹⁷ Haggard and Cochran⁹⁸ and Bahadori *et al.*,¹⁰⁴ predictions from Klajn and Oppenheim¹⁰² and Edelman *et al.*,⁹⁵ plot from Bahadori *et al.*,¹⁰⁴

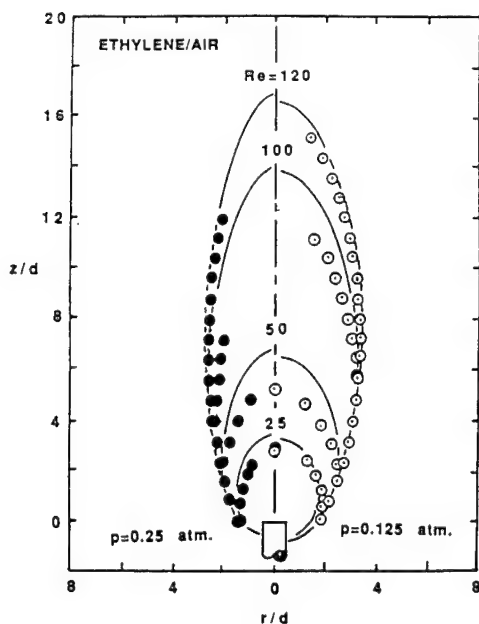


FIG. 21. Predicted and measured shapes of weakly-buoyant ethylene-air laminar jet flames at various burner exit Reynolds numbers. From Mortazavi *et al.*,¹⁰⁸

tions of the boundary layer and the negligible radiative heat loss approximations.¹⁰⁸⁻¹¹⁰ The approach of Mortazavi *et al.*¹⁰⁸ and Köylü *et al.*¹⁰⁹ attempts to address the difficulties of soot-containing flames by using the conserved-scalar formalism in conjunc-

tion with the state relationships discussed in connection with Fig. 17. The flames were taken to be axisymmetric and steady, while radiative heat losses were treated in an approximate way by assuming that they were a fixed fraction of the chemical energy release, similar to treatments used with some success for turbulent diffusion flames.⁸⁷ Measurements to evaluate the predictions were carried out for weakly-buoyant laminar jet flames at ng. This involved measurements at reduced pressures, exploiting the fact that effects of buoyancy for laminar diffusion flames scale like $(p/p_0)^2 g$, where p and p_0 are the test pressure and normal atmospheric pressure. Thus, flames at pressures of $O(0.1 \text{ atm.})$ have buoyancy effects of $O(0.01 g)$ and can be made weakly buoyant by controlling the jet exit velocity if they are not too long. Predicted and measured flame shapes using a 3 mm diameter burner for ethylene-air flames are illustrated in Fig. 21; results for acetylene-air flames were similar. The measurements denote the position of the blue portion of the flame and are terminated when yellow soot luminosity obscures the blue flame sheet; the predictions denote the corresponding locus of the stoichiometric mixture fraction. Predictions include effects of buoyancy, but they are nearly negligible for these conditions; this behavior is evident from the small change of flame shape when the pressure is halved. The predictions are reasonably good, including the trends of the flame shape with varying Reynolds number and the attachment of the flames below the burner exit. In particular, capabilities to predict features near the burner exit and the shapes of low Reynolds number flames demonstrate the value of allowing for streamwise diffusion rather than using the boundary layer approximation.

The evaluation of the approximate approach for computing the properties of soot-containing diffusion flames was continued by Köylü *et al.*¹⁰⁹ In this study, distributions of velocities, mixture fractions and the concentrations of major gas species were considered for weakly-buoyant laminar jet diffusion flames. Typical predictions and measurements are illustrated in Fig. 22 for an acetylene-air flame at 0.250 atm. The state relationships used for these predictions were drawn from results at atmospheric pressure from Gore and Faeth,⁸⁵ nevertheless, concentration measurements in the low-pressure flames indicated the small effect of pressure on the state relationships over the range of interest. The weakly-buoyant nature of the flame is indicated by the rapid decay of velocity near the burner exit; velocities for more buoyant atmospheric pressure conditions would begin to increase almost immediately. Nevertheless, a gradual increase of velocity is still observed at larger distances from the burner exit which is caused by buoyancy—highlighting problems of ng observations of flames of reasonable size having negligible effects of buoyancy discussed in Section 2. In any event, the approach of Refs

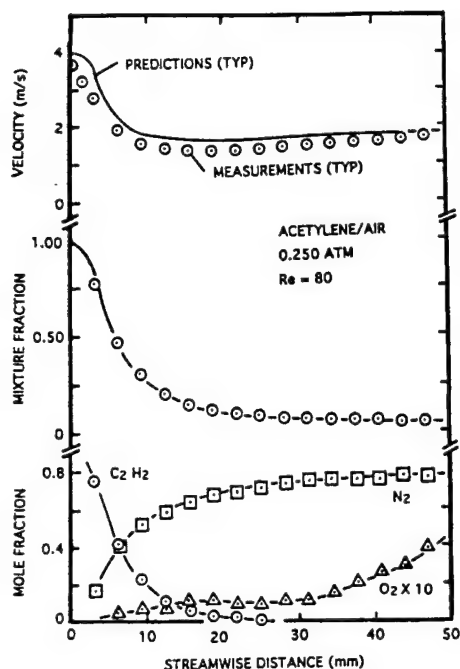


FIG. 22. Predicted and measured velocities, mixture fractions and mole fractions of acetylene, nitrogen and oxygen along the axis of a weakly-buoyant acetylene-air laminar jet diffusion flame at 0.250 atm. From Köylü *et al.*¹⁰⁹

108 and 109 handles the mild effect of buoyancy reasonably well, and predictions are in reasonably good agreement with measurements. Evaluations of temperature distributions in the fuel-lean portions of the test flames were also reasonably satisfactory.^{108,109}

The results of Refs 108 and 109 are promising; however, predictions of this type still have considerable uncertainties. First, the flame shape is sensitive to radiative heat losses that are crudely modeled; this sensitivity to radiation would present greater difficulties for atmospheric pressure ethylene and acetylene flames that have much higher soot concentrations, and thus much larger continuum radiation heat losses from soot. Second, the variation of soot concentrations with pressure raises questions about the relevance of state relationships measured at atmospheric pressure to these low-pressure conditions, although pressure effects on state relationships have proven to be small for ethylene and acetylene-air flames for pressures in the range 0.125–1.000 atm.,¹⁰⁹ and the existence of generalized state relationships for hydrocarbons having a wide range of sooting tendencies provides some justification for this procedure.⁸⁶ Finally, additional structure measurements for a wider range of conditions, including effects of coflow, are needed for a definitive evaluation of the predictions.

Interest in reliable predictions of the structure of nonbuoyant laminar jet flames goes beyond the general objective of understanding this fundamental flame configuration. For example, laminar jet diffu-

sion flames at μg provide a useful experimental environment, capable of addressing a wide range of pressures, for studies of fuel decomposition, soot processes and pollutant production in diffusion flames. However, diagnostics available to characterize the flame environment at μg will be limited in comparison with typical laboratory conditions at ng for some time to come. As a result, reliable methods to predict the main features of flame structure—concentrations of major species, temperatures and velocities—are needed in order to supplement limited measurements to define the flame environment. Thus, additional experimental and theoretical work for nonbuoyant flames is needed to establish reliable prediction methods. In particular, similar to premixed combustion at μg , radiation plays a strong role in the properties of nonbuoyant jet diffusion flames and will require greater attention than in the past to achieve this goal.

4.3. Flame Stability

Gaseous diffusion flames involve mixing between fuel-rich and oxidant-rich streams, with a flame zone appearing in the mixing region between the two streams. Various configurations of these streams are used in practice, however, they are canonically similar to the jet diffusion flame configuration. Thus, jet diffusion flames will be used to simplify the discussion of flame stability issues in the following. The main problems of stability for nonpremixed flames involve maintaining the flame attached at the jet exit, local extinction of the flame sheet due to excessive stretch, and transition to turbulence.

The stability of flame attachment determines whether the flame remains attached near the jet exit, whether the flame is lifted or separated to some distance from the jet exit, or whether the flame blows off entirely and is extinguished. In the presence of buoyancy at ng, or for strong forced-convection conditions, the mechanism of attachment is considered to involve local premixing of fuel and oxidant which provides a small premixed flame region. This behavior comes about because the flame attachment point is generally separated from the burner exit, providing a region where the fuel- and oxidant-rich streams can mix without reaction to form a flammable gas mixture. Propagation of the resulting premixed flame is then stabilized by local gas velocities, which must be low enough so that the flame can maintain a fixed position. Additionally, for attached flames, where the premixed flame is close to the burner exit, heat loss to the burner provides an additional stabilization mechanism, similar to the behavior of fully premixed flames stabilized on flat-flame burners. Finally, the premixed flame region can be viewed as providing a continuing ignition source that anchors the position of the rest of the flame.

This traditional view of flame attachment could very well be modified for low Reynolds number flames at μg . For example, observations of laminar jet diffusion flames in a stagnant environment at μg show that the points of flame attachment advance below the burner exit,⁸⁶⁻⁸⁸ similar to the predicted flame shape plots for weakly-buoyant flames illustrated in Fig. 21. In the complete absence of buoyancy, and for negligible jet momentum, diffusion in all directions is equally probable so that the flame shape would become spherical and the attachment points would move farther down the tube. In this case, conventional attachment is no longer required, any more than it is required for the envelope flame around a motionless burning drop at μg , and the attachment points would simply be areas where quenching occurs due to the relatively cool burner surface. Clearly, the evolution between the spherical flame and the more conventional attachment of flames having large velocities involves progressive increases of stabilization due to premixing effects, and can offer useful insights about the mechanism of attachment.

The attachment process is at least two-dimensional; therefore, attachment cannot be described by boundary-layer approximations because streamwise transport acts to fix the attachment point, and attachment involves complications of evolution from a premixed to a diffusion flame. In view of these complexities, current understanding of attachment and extinction is very limited—largely consisting of empirical correlations of measurements based on phenomenological theories.^{41,42} Experimentation at μg , however, should advance our understanding of processes of attachment and extinction because the absence of buoyant velocities allows the stretch of the premixed region to be controlled by jet exit velocities, and modified systematically by controlled coflow velocities, while capabilities to operate at low jet exit Reynolds numbers expands the premixed flame region so that adequate spatial resolution for measurements can be provided. Models allowing for streamwise diffusion and finite-rate kinetics (for example, Ref. 110) can potentially treat the attachment regions, but corresponding measurements have not been addressed as yet.

Problems of local extinction of diffusion flame surfaces are important as potential sources of pollutants in turbulent flames.⁹¹ Thus, stretched laminar flame surfaces are receiving a great deal of attention using experiments at ng. Companion tests at μg would also be helpful, due to the potential for improved spatial resolution and the need to address the low stretch rates of spacecraft environments for fire safety concerns. However, measurements at μg will require development of instrumentation comparable with methods available at ng, as well as extended access to spacecraft μg facilities. Thus, instrument and facility development must precede significant contributions of μg experi-

ments to a better understanding of local flame quenching.

Issues of transition from laminar to turbulent flames are long standing due to problems of buoyant disturbances in the transition region. Bahadori *et al.*¹¹¹ report initial results concerning transition to turbulence in round jet diffusion flames at μg , finding significantly different behavior in the transition region for nonbuoyant and buoyant conditions. In particular, large-scale slow-moving wrinkled flame structures are observed near the flame tip at μg , rather than the brush-like flame tip seen at ng. Additionally, transition to a turbulent flame at μg is characterized by the appearance of intermittent disturbances that are generated near the flame base and convected downstream, in contrast with ng flames where disturbances are first observed at the flame tip. These fundamental differences are probably related to the different velocity distributions in nonbuoyant and buoyant laminar diffusion flames. In particular, velocities decrease with increasing streamwise distance for nonbuoyant flames (Fig. 22), but increase with increasing distance for buoyant flames. This behavior would tend to move regions of high velocity, which are most unstable to transition, downstream toward the flame tip at ng in agreement with the observed behavior. However, the actual mechanism for different transition behavior at μg and ng still is unknown. Finally, flame lengths increase monotonically with increasing burner exit Reynolds number at μg , rather than the abrupt reduction of the flame length at transition seen at ng. Clearly, existing μg facilities are capable of addressing issues of transition to turbulence in flames and they deserve consideration due to the importance of turbulent flames to practical applications.

4.4. Soot Processes

Understanding soot properties in diffusion flames is important because soot affects the performance of propulsion systems and furnaces, the hazards of unwanted fires, and the pollutant emissions from combustion processes. For example, continuum radiation from soot is the main heat load to the combustor components of propulsion systems and controls their durability and life, while it also serves as the main heat transfer mechanism in many furnace designs.¹¹² Similarly, continuum radiation from soot is the dominant mechanism for the growth and spread of unwanted fires, particularly for spacecraft conditions where effects of buoyancy are absent, while soot-containing clouds emitted from these flames obscure fire-fighting efforts.^{87,113} Finally, black soot-containing exhaust plumes and the carbon monoxide emissions intrinsically associated with soot emissions^{114,115} represent objectionable pollutants and are also the main source of fatalities in unwanted fires.¹¹⁶ Thus, soot properties within diffusion flames

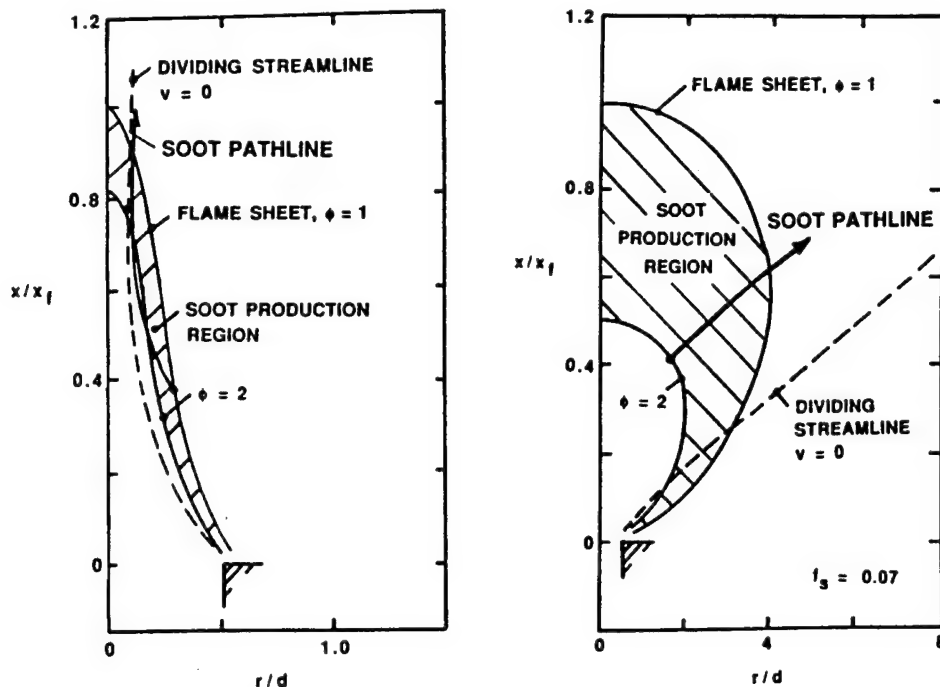


FIG. 23. Schematics of soot formation and oxidation regions for laminar jet diffusion flames at ng and μ g.

are unusually significant for practical applications, and they represent a major unresolved problem of combustion science in spite of significant efforts in the past.^{117,118}

Similar to other flame phenomena, while soot processes in turbulent flames are of greatest practical interest, the direct study of soot in turbulent flames is not feasible using either existing or anticipated technologies. In particular, the unsteadiness and distortion of turbulent flames limit available residence times and spatial resolution in regions where soot properties are most important. Thus, laminar diffusion flames are generally used as more tractable model flame systems to study soot processes relevant to turbulent diffusion flames, justified by the known similarities between the gas-phase processes in these two-flame systems.^{86,90} This is appropriate only if the buoyancy in laminar diffusion flames at the ambient pressures of interest, generally atmospheric pressure and greater, does not directly affect soot processes. Unfortunately, this is not the case because soot particles are too large to diffuse like gas molecules and are primarily transported by local convective flows, modified somewhat by effects of thermophoresis.

The problems associated with using buoyant laminar flames to study soot processes can be seen by comparing the configurations of the soot production regions in buoyant and nonbuoyant laminar jet diffusion flames as illustrated in Fig. 23. Soot production regions occur for fuel-equivalence ratios roughly in the range $\phi = 1-2$,¹¹⁸ which is marked on the plots.

The locus of points where the radial velocity, $v = 0$, which approximates the dividing streamline, is also shown on the plots, along with some typical soot pathlines in the flow. Soot particles follow the flow along streamlines, ignoring effects of thermophoresis; therefore, they move toward the dividing streamline in the radial direction. For buoyant flows, the dividing streamline is inside the soot production region except near the flame tip. Thus, except for a small region near the axis, soot particles form near the flame sheet and then move radially *inward* toward cooler and less reactive regions before finally reversing direction, in mixture fraction space, and eventually crossing the flame sheet close to the flame tip in a narrow soot layer near the dividing streamline. However, the process is completely different for nonbuoyant jet diffusion flames because the dividing streamline crosses the flame sheet close to the burner exit; as a result, soot begins to form near the cooler core of the flame and is drawn directly toward and through the flame sheet for most of the flame. Another important difference between buoyant and nonbuoyant diffusion flames involves the variation of velocities along soot pathlines. For buoyant flames, velocities progressively increase with increasing height above the burner exit, for example, similar to the discussion leading to Eq. (3), $(\Delta \rho g z / \rho)^{1/2}$ is roughly the scale of streamwise velocities. Thus, residence times near the upper end of the soot reaction zone are shortest for buoyant flames, which reduces the effectiveness of soot oxidation processes in comparison with soot production processes nearer to the

burner exit. In contrast, streamwise velocities are roughly inversely proportional to the distance from the burner exit for nonbuoyant flames, as discussed in connection with Fig. 22.^{108,109} This tends to increase the effectiveness of soot oxidation processes relative to soot growth processes for nonbuoyant flames in comparison with buoyant flames. Finally, as will be discussed later, residence times in nonbuoyant flames are significantly longer than for buoyant flames of comparable size, providing longer absolute times for soot nucleation, growth and oxidation. Thus, any resemblance between soot processes within nonbuoyant and buoyant laminar diffusion flames clearly is fortuitous. Additionally, local effects of buoyancy are generally insignificant for turbulent flames; thus, the largely unstudied nonbuoyant laminar jet diffusion flame configuration provides a better simulation of soot processes relevant to the turbulent flames of practical interest than buoyant laminar jet diffusion flames.

Aside from their practical relevance, nonbuoyant jet diffusion flames at μg also provide improved spatial resolution for studying soot processes. This can be seen from the results illustrated in Fig. 23. In particular, the flame surface and the dividing streamline are close to one another for buoyant flames so that soot collects in a narrow layer. Furthermore, soot oxidation is confined to a narrow region where the soot layer crosses the flame tip for buoyant flames, because effects of strain and relatively high velocities in the flow near the flame tip introduce large radial gradients as well as rapid quenching of reactions in the soot layer. In contrast, both soot formation and oxidation are spread along most of the flame surface for nonbuoyant conditions, and these regions are broad with relatively low velocities for the low Reynolds number conditions that are accessible at μg . In particular, the breadth of the soot-containing region is evident from the significant flame widths seen in Fig. 21.

Flexibility to control overall flame residence times, defined as the time required for a parcel of fluid to convect from the burner exit to the flame tip, is another advantage of nonbuoyant jet diffusion flames for studying soot processes. For example, flame residence times for buoyant laminar diffusion flames are primarily controlled by effects of buoyancy and are relatively insensitive to burner diameter and initial gas velocity variations, with the residence time roughly proportional to the square root of the flame height.^{114,115} This behavior implies a variation of residence times of approximately 3:1 over the range of typical laminar smoke point flame lengths for buoyant flames, for example, 20–200 mm, at atmospheric pressure.^{117–120} In contrast, residence times tend to be proportional to burner diameter and inversely proportional to burner exit velocity for nonbuoyant laminar jet diffusion flames having dimensions typical of laminar smoke point conditions at atmospheric pressure, that is, burner exit Reynolds

numbers of $O(100)$ and ratios of flame length to burner exit diameter less than 50.¹⁰⁸ Based on simple constant property analyses using the boundary layer approximations, the effect of diameter is expected; however, the effect of burner exit velocity is surprising because simplified analyses show that residence times are independent of the burner flow rate.¹¹⁷ The difference between the two results is caused by effects of streamwise diffusion and variable properties; actually, due to the large property variations within flames, rather large burner exit Reynolds numbers, $O(1000)$, are required before the boundary layer approximations are appropriate for laminar jet diffusion flames.¹⁰⁸ Thus, it is a relatively simple matter to vary the residence times of nonbuoyant jet diffusion flames by orders of magnitude through variations of burner exit diameters and velocities for the range of conditions normally considered during studies of soot processes.

Recent studies of soot processes in laminar jet diffusion flames at μg have confirmed the anticipated differences between soot processes at ng and μg conditions, and have demonstrated the advantages of experiments at μg . A portion of these studies was devoted to measurements of the laminar smoke point flame length, which is defined as the length of the luminous portion of the flame at the condition where soot just begins to be emitted from the flame, that is, all longer flames which have larger fuel flow rates will be soot emitting. The laminar smoke point flame lengths of buoyant jet diffusion flames at ng are a valuable global measure of the sooting tendencies of fuels, because these lengths, and the corresponding residence times for a particular length, are relatively independent of burner exit conditions.^{114–120} It was suggested earlier¹¹⁷ that laminar smoke point flame lengths might not be observed for nonbuoyant laminar jet flames because simplified boundary layer analysis indicated that their residence times were independent of burner flow rate, or flame length for a particular burner, as discussed earlier. However, laminar smoke point flame lengths were recently observed for nonbuoyant flames at μg which provide a useful contrast to existing observations at ng.¹²¹ A portion of the results is summarized in Table 1, which provides laminar smoke point luminosity lengths and residence times for ethylene and propane–air flames at atmospheric pressure. Luminosity lengths are somewhat longer than flame lengths because the soot oxidation region is relatively large; however, the ratios between flame and luminosity lengths are comparable at ng and μg , ca. 0.6.¹²¹ On the other hand, the basis for residence times differ somewhat because they were computed using the approach of Mortazavi *et al.*¹⁰⁸ at μg , while they were measured as the time required for flame luminosity to disappear after abruptly ending the burner flow at ng.¹⁹ It is unlikely, however, that the different definitions of residence times significantly affect the following observations.

TABLE 1. Laminar smoke point properties for nonbuoyant and buoyant diffusion flames

Buoyancy condition	Burner diameter (mm)	Luminosity length (mm)	Residence time (ms)
<i>Ethylene-air flames</i>			
Nonbuoyant (μg)	1.6	32	180
Nonbuoyant (μg)	2.7	25	280
Nonbuoyant (μg)	5.9	24	750
Buoyant (ng)	14.3	135	41
<i>Propane-air flames</i>			
Nonbuoyant (μg)	1.6	42	230
Nonbuoyant (μg)	2.7	38	420
Nonbuoyant (μg)	5.9	42	1310
Buoyant (ng)	14.3	169	48

Nonbuoyant and buoyant round jet flames in air at 298 ± 3 K and 1 atm. Nonbuoyant results from Sunderland *et al.*¹²¹ with residence times estimated from Mortazavi *et al.*¹⁰⁸ Buoyant flames from Sivathanu and Faeth¹¹⁹ with measured residence times.

There are several interesting features about the comparison between laminar smoke point luminosity lengths and residence times at μg and ng summarized in Table 1. First, the nonbuoyant flames exhibit laminar smoke point luminosity lengths, in contrast with the implications of simplified theories of flame structure,¹¹⁷ because their residence times vary with flame length as discussed earlier. Second, varying the burner exit diameter at μg has little effect on the laminar smoke point luminosity length. While this behavior is superficially similar to buoyant flames, the mechanism must be different. In particular, the residence times increase substantially over the range of burner diameters at μg listed in Table 1; however, similar variations of burner diameter would have little effect on residence times at ng.¹¹⁷ Additionally, the smoke point luminosity lengths are roughly four times shorter at μg than ng. Finally, residence times at the laminar smoke point are appreciably larger at μg than at ng—180–1310 ms in comparison with 41–48 ms. About the only feature that is similar for results at μg and ng is the tendency for luminosity lengths and residence times to be somewhat larger for propane-air than ethylene-air flames. The different soot paths and the different relative residence times and sizes of the soot formation and oxidation regions for nonbuoyant and buoyant jet diffusion flames undoubtedly contribute to this behavior. However, the specific mechanisms are neither adequately quantified nor understood. Furthermore, the greater propensity to soot at μg than ng is problematical for fire safety in spacecraft. Clearly, much remains to be done to establish the behavior of smoke point properties at μg , but it is already evident that experiments at μg provide a valuable new perspective for gaining a better fundamental understanding of the global soot properties of diffusion flames, as well as information needed to address fire safety concerns for spacecraft.

Other experiments carried out at μg further con-

firm fundamental differences between soot processes in nonbuoyant and buoyant jet diffusion flames.^{93–107} These experiments involve jet diffusion flames for a variety of hydrocarbon fuels in still air with the measurements limited to flame photographs. The results were generally obtained using drop towers where short available test times preclude reaching fully-developed flame conditions; however, the results are still qualitatively useful.¹⁰⁶ It was observed that μg conditions yield enhanced sooting, which was attributed to the longer residence times of μg flames than ng flames of comparable lengths.¹⁰³ The region where soot was present was much wider at μg than at ng, with the luminous region due to continuum radiation from soot having a blunt tip, called tip-opening behavior, rather than the familiar conical tip of buoyant flames such as candle flames. Furthermore, the luminosity near the tip of nonbuoyant flames progresses from yellow to red to dull red with increasing distance from the burner exit, near the end of the visible portion of the flame. This behavior can be attributed to the absence of buoyancy as discussed in connection with Fig. 23. For nonbuoyant and weakly-buoyant conditions, soot crosses the flame sheet over a broad region and passes into an extended soot oxidation region where gas temperatures progressively decrease; this yields a blunt appearance to the luminosity emitted from the soot-containing region with the color shifting toward the infrared as the mixture cools. In contrast, the soot layers in buoyant flames at ng are confined to a region near the axis as the flame tip is approached, which yields a conical tip because the oxidation region is rapidly quenched.

Recent measurements of soot concentrations in weakly-buoyant laminar jet diffusion flames at low pressures help to quantify the structure of the soot-containing region when the effects of buoyancy are small.¹⁰⁸ These experiments were feasible because even though soot concentrations are reduced at low

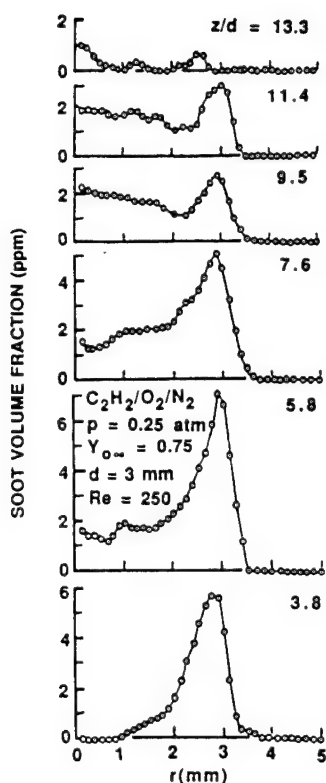


FIG. 24. Soot volume fraction distributions in a weakly-buoyant laminar jet diffusion flame (acetylene/ O_2/N_2 mixture at 0.125 atm, and an ambient oxygen mass fraction of 0.75). From Mortazavi *et al.*¹⁰⁸

pressures,^{122,123} longer residence times in nonbuoyant flames help to compensate for this reduction. A typical example of these results is illustrated in Fig. 24, which involves an acetylene flame issuing from a 3 mm diameter burner having a burner exit Reynolds number of 250, an ambient oxygen mass fraction of 0.75 and an ambient pressure of 0.25 atm. The soot volume fraction is plotted as a function of radial distance at various distances from the burner exit. An annular soot layer is evident near the burner exit, corresponding to the region that is sufficiently heated for soot to form with possibly some additional resistance to the soot crossing the flame sheet due to effects of thermophoresis. However, the soot layer becomes less prominent as soot begins to form along the axis of the flow near the tip of the flame (ca. $x/d = 9.35$, based on the stoichiometric mixture fraction). Soot oxidation extends to the axis beyond the flame tip, where soot concentrations become relatively uniform. Finally, the soot is completely oxidized at a nearly constant distance from the burner exit in this nonsooting flame. The large width of the soot-containing region, and its rather uniform disappearance at all radial locations, helps to explain the blunt luminous zone (or tip-opening process) observed for jet diffusion flames at μg . Finally, the breadth of the soot-containing region and the large

size of the soot-oxidation region clearly provide excellent spatial resolution for studying soot processes.

Behavior similar to Fig. 24 at all pressures at μg , coupled with significant flexibility to vary flame residence times, provides unprecedented opportunities for significant progress towards an understanding of soot processes in diffusion flames. The differences in soot behavior at μg and ng have important implications for spacecraft fire safety because continuum radiation from soot significantly affects fire growth, flame spread and burning rates,⁸⁷ providing an additional incentive to study soot processes in nonbuoyant diffusion flames.

5. HETEROGENEOUS PREMIXED FLAMES

5.1. Introduction

Aside from the limited case of double-base solid propellant combustion, denoting a class of combustion phenomena as heterogeneous premixed flames is an oxymoron because these flames normally involve nonpremixed combustion on small scales, that is, individual dispersed phases (particles or droplets) are generally dominated by either fuel-like or oxidizer-like reactants. In keeping with normal usage, however, we shall consider a variety of flame phenomena under the heading of heterogeneous premixed flames, where gross mixing between the fuel-rich and oxidizer-rich phases has already occurred before flame propagation begins. This includes flame propagation in suspensions, smoldering and materials synthesis in flames.

5.2. Flame Propagation in Suspensions

The propagation and extinction of flames in particle and droplet suspensions are relevant to the combustion of sprays and coal dusts, and to accidental explosions in mine galleries and grain elevators. However, it has proven to be far more difficult to develop reliable information concerning such important fundamental combustion parameters as the minimum ignition energy, flame propagation speeds, flammability limits, and quenching distances in particle and droplet suspensions than in homogeneous gas mixtures. The primary reason for the difficulties is the inability to produce a uniform suspension due to particle settling. This problem is caused by two factors, namely particle settling during combustion, and particle settling during the preparation of a combustible heterogeneous mixture.¹²⁴⁻¹³⁰

First consider the effects of particle settling during combustion. The particles of interest to combustion have sizes which typically vary from 10–1000 μm . This implies terminal velocities within the range

10–100 cm/s. This range, however, basically brackets the rate of flame propagation in particle suspensions so that settling can seriously modify the flame behavior. For example, for a downward-propagating flame, rapid settling may make the suspension too dilute to support combustion. In contrast, an upward-propagating flame may also fail to propagate because the dense particle cloud can either make the heterogeneous mixture too rich for flame propagation and/or can absorb too much heat from the flame and cause extinction. By the same reasoning, there may also exist situations in which settling can make an initially nonflammable suspension combustible. Finally, even if a flame can successfully propagate through the suspension, its motion will be highly transient due to the stratified particle concentration.

The second factor concerns the creation of particle or droplet clouds. Effective methods to mix particles and gases frequently produce strong turbulent motion within the gas (see Kumar *et al.*¹³⁰ and references therein). Since it is desirable to initiate the experiment only after the turbulent motion has decayed, substantial settling can take place during this decay period. Practical operational compromises imply that measurements at μg involve combined disturbances of turbulence and settling except for the very smallest particulates or droplets.¹³⁰ In view of these considerations, the need for long duration μg experiments in space is warranted for this class of phenomena. These experiments will provide valuable data for comparison with existing theories based on uniform suspensions (see Seshadri *et al.*¹³¹ and references therein), and will serve as new standards for the evaluation of fire hazards in particle suspensions for both Earth- and space-based applications.

Earlier μg studies on flame propagation in suspensions in a tube have focused on a so-called 'chattering flame' phenomenon.^{125–128} Specifically, the structure of near-stoichiometric flames appears to be spatially discontinuous in a rib-like manner, while it is continuous for rich mixtures. It was speculated that the chattering flame propagation was associated with a Kundt's tube-type phenomenon,¹³² which causes the unburnt mixture to segregate into alternate fuel-rich and fuel-lean layers. Upon ignition, the flame would then propagate in a pulsating mode as the layers of particles are ignited in turn through radiation. More recent experiments, however, cast doubt on this mechanism because segregated layers in the unburnt mixture were not observed.¹³³ It is now believed that the chattering nature is simply caused by the flame-acoustic interaction for flame propagation in a tube.

Theoretical studies for flame propagation in suspensions have been performed for both the steady mode of flame propagation,¹³¹ and for the acoustic instability characteristics in the tube.^{133,134} In all these analyses the particles are assumed to completely gasify in the preheat zone so that the reaction zone is equivalent to a premixed flame. Results presented

for acoustic instability characteristics in Ref. 133 show that the growth of the instability is damped by the particle velocity lag, as is well known, with the damping being stronger for the higher loading cases of stoichiometrically-rich mixtures.

Concerning future research, it seems that if the chattering flame phenomenon is strictly a flame-acoustic interaction problem peculiar to the use of the tube as the flame propagation chamber, then investigators can simply use either a homogeneous gaseous mixture for studies on acoustic instability alone, or a gas mixture with inert particle seeding to allow for viscous damping. By the same reasoning, studies of flame propagation in fuel particle suspensions should use a combustion chamber which would not produce such instabilities. In terms of theoretical work, active particle gasification and burning should constitute part of the reaction zone structure, similar to 'spray flame' analysis,²⁴ so that the heterogeneous nature of the flame structure is completely incorporated.

5.3. Smoldering

Accidental fires and explosions are frequently preceded by smoldering, which is the slow heating and pyrolysis of organic materials with the release of combustible vapor and particles. Without buoyancy for dispersion, the pyrolyzate will stagnantly accumulate so that a flammable concentration can be easily reached. In addition, safe toxicity levels in the environment could be exceeded due to accumulation of these preflame pyrolyzates. Therefore, the formation and dispersion of pyrolyzates for smoldering in μg environments needs to be understood, primarily motivated by spacecraft fire safety concerns.^{135,136}

Smoldering frequently takes place in the interior of a porous material through the creeping propagation of a smoldering front supported by the exothermic, heterogeneous reaction that takes place at the surface of the pores. Radiative transport may become important because heat conduction can be relatively ineffective in a porous medium. Furthermore, because the porous material inhibits convective and diffusive transport to the smoldering front, oxygen supply is frequently the rate limiting factor. Smolder front propagation has been studied for cigarettes, polyurethane foams, wood, wood products, coal matrixes in underground coal gasification and fires involving coal heaps.¹³⁵

The influence of gravity on smoldering is primarily through the mechanism and rate of air supply, with past considerations generally limited to downward and upward propagation (see Refs 136–138 and references therein). For example, in a downward-propagation mode, the supply of oxidizer is augmented by the buoyancy-induced upwardly-rising air. Since both the fuel and oxidizer are supplied upstream of the front, the spreading mode is similar to a premixed

flame and is termed cocurrent spread. On the other hand, for an upwardly-propagating flame, fuel and buoyancy-induced oxidizer are supplied from opposite sides of the flame. Then the spreading mode is similar to a nonpremixed flame and is termed counter-current spread. The above situations are analogous to those of flame propagation in the vertical tube, considered previously, except that the flow is in a porous medium. The spreading of the smoldering front can also take place through natural convection boundary-layer flows over nonporous surfaces for the analogous cocurrent and counter-current situations. It is clear that, in spite of the small pore size of the smoldering material, buoyancy plays an important role in the rates of flame propagation and spreading in these situations. In the absence of buoyancy, the rates of oxidizer supply, and thereby front propagation, are expected to be considerably reduced. Detection of smoldering occurrence will also be more difficult when buoyancy is absent, leading to greater danger of transition to flaming combustion.

The μg environment also provides an opportunity for systematic study of smoldering processes that cannot be carried out at ng. For example, detailed measurements of the locally heterogeneous combustion processes of smoldering requires reasonable spatial resolution. However, this is difficult at ng because gas passage sizes must be small to prevent effects of natural convection flows that are not relevant to practical smoldering processes (see Section 2 for a discussion of these limitations). This difficulty is circumvented at μg because the size of both the fuel elements and the gas passages can be scaled up so that conventional temperature and gas sampling probes can be used to determine flame structure. Unfortunately, smoldering processes are normally slow, and become even slower upon scale-up. Thus, ground-based μg facilities, that have test times in the range 1–100 s, do not offer much potential for studies of smoldering. However, progress in this area should be rapid when spacecraft environments become available for smoldering experiments.

5.4. Materials Synthesis

Materials synthesis in flames is a growing field due to potential applications for a wide range of novel and useful products. There are three general classes of materials synthesis in flames: film deposition, condensation of nanoparticles and self-propagating high-temperature synthesis (SHS). Flame deposition of films has a long history and is commonly used to produce coatings of refractory materials, including recent interest in the production of industrial diamonds. Significant effects of buoyancy on deposition processes are normally not a factor, however, because relatively high flow velocities are used in order to minimize the boundary layer thickness along surfaces so that high deposition rates can be achieved.

Thus, deposition will not be considered in the following discussion. However, materials synthesis by condensation of nanoparticles and SHS will be considered because they are related to flame propagation in suspensions and smoldering so that μg environments can impact these processes as discussed in Sections 5.2 and 5.3.

A common method of materials synthesis in flames involves condensing the desired product in the post-flame region, somewhat analogous to the way that soot is formed in premixed flames. This tactic can be used to produce highly purified refractory materials like silicon dioxide or technological fumes like carbon black and fullerenes.¹³⁹ In a general way, this is the inverse process of flame propagation in suspensions and it involves similar limitations. First of all, the growth of dispersed-phase elements is limited by settling and available flame residence times at ng. At μg conditions, both these restrictions are modified, thereby allowing for the production of larger dispersed-phase elements, providing a range of properties of the materials that is beyond current capabilities, and reducing problems of phase separation (or product collection). However, the properties of these processes are not understood at μg ; in particular, it is likely that available residence times will ultimately be limited by radiation effects, rather than convection effects which control residence times at ng. Thus, the kinds of products that can be made, and the structure and stability of the flame environments used for their production, will require extensive experiments at μg .

SHS is related to thermite processes that have a long history, and is receiving new interest due to capabilities to produce novel and valuable products (see Refs 140–142 for a summary of recent work in this field). SHS involves mixing and compacting controlled amounts of solid phases of reactant materials and igniting the mixture, with the final product produced upon flame propagation through the mixture. A desired shape of the product also can be achieved by appropriate shaping of the compacted reactant. There are two main classes of SHS processes: (i) solid–fluid and (ii) solid–solid. Solid–fluid processes involve one reactant as a solid matrix that is submerged in a second reactant that is fluid. An example is the suspension of titanium particles in liquid nitrogen to produce titanium nitride.¹⁴³ Solid–solid processes involve mixtures of solid phases, typically of a metal and a nonmetal, with the surrounding gas playing a minor role, or with operation even in a vacuum, yielding a unique class of combustion phenomena. Solid–solid SHS processes are sometimes called gasless reactions because both reactants and products are solids; however, one of the reactants frequently melts or gasifies upon passage of the flamefront so that the resulting combustion process could resemble heterogeneous flame propagation in sprays and dusts.¹⁴⁴ Including both classes, there is a rich variety of SHS processes of interest, and the

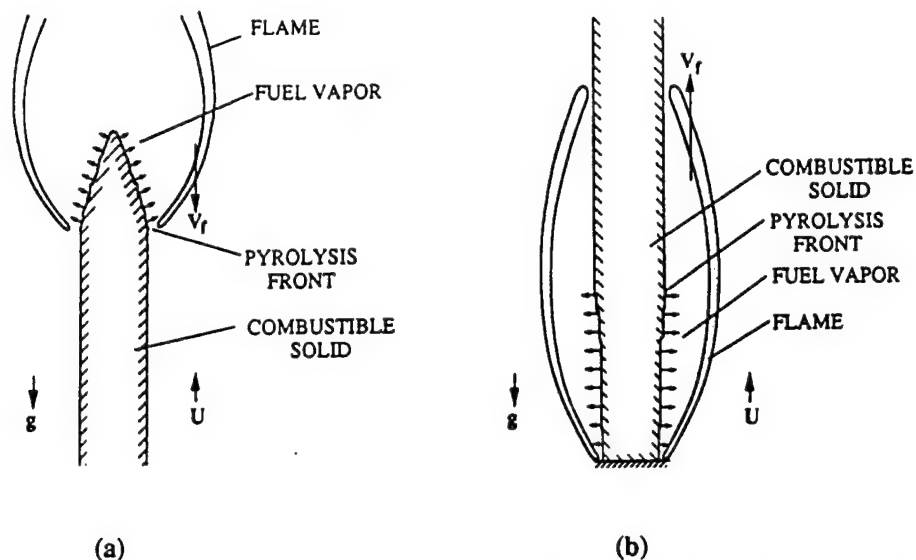


FIG. 25. Schematic of (a) a downward-propagating opposed flow flame, and (b) an upward-propagating concurrent flame.

mechanisms of SHS are not well understood. In fact, most past work has been limited to general thermodynamic considerations to define conditions needed to achieve SHS having appropriate properties, and studies of the mechanical properties of the product for various reactant properties.¹⁴⁰⁻¹⁴² Thus, there is substantial scope for fundamental experimental studies of SHS, with the advantages of experimentation at μg being very similar to those already discussed in connection with heterogeneous premixed flames. There are some additional distinct advantages of conducting the SHS process at μg . For example, for solid-solid synthesis in which the flame speed could be low (say 0.1 cm/s), and one of the solids melts upstream of the reaction front, the melt could drip due to gravity and consequently modify the unreacted mixture composition. Furthermore, for liquid-solid synthesis involving a suspension, the problem of settling could be circumvented at μg in the same manner as for gas-particle suspensions discussed in Section 5.2, provided a uniform dispersion can be initially achieved.¹⁴³

6. HETEROGENEOUS NONPREMIXED FLAMES

6.1. Introduction

As noted earlier, the combustion process of heterogeneous flames generally involves nonpremixed combustion between a gaseous phase and a solid or liquid phase. Therefore, heterogeneous nonpremixed flames include many classical problems that have had significant attention in the past. In the following we shall consider two main classes of heterogeneous nonpremixed combustion, namely, flame spread along solid and liquid surfaces and the combustion

of droplets. The corresponding problems of the combustion of solid particles has attracted surprisingly little attention at μg conditions, and will not be considered.

6.2. Flame Spreading

Flame spreading over fuel surfaces is a crucial process in fires both on Earth and in space. The relevant issues are understanding the mechanisms governing spreading, determining the spreading rate as a function of the fuel properties and environmental parameters, and identifying the limiting situations when spreading is not possible.

Flame spreading can be classified into two broad categories,¹⁴⁶ namely, opposed-flow spread (Fig. 25a), in which the spreading direction is opposite to that of the external flow, and concurrent-flow spread (Fig. 25b), in which the spreading direction is the same as that of the external flow. For a vertically-oriented fuel surface in the presence of gravity, the flame induces a buoyant flow and these two spreading modes correspond to downward and upward flame spread, respectively. In concurrent spread, the hot combustion products form a plume that bathes the unburnt fuel, leading to rapid rates of flame spreading. If the fuel plate is also thermally thick,¹⁴⁶ then spreading is generally accelerated as a greater depth of the solid fuel becomes heated. For a thermally thin fuel, a steady rate of spread may be possible when complete fuel burnout can be achieved near the leading edge. Opposed spread is slower in general and a steady state can be achieved more readily. Due to its relative experimental and theoretical simplicity, most research has been conducted on opposed spread of laminar flames over thin fuels.

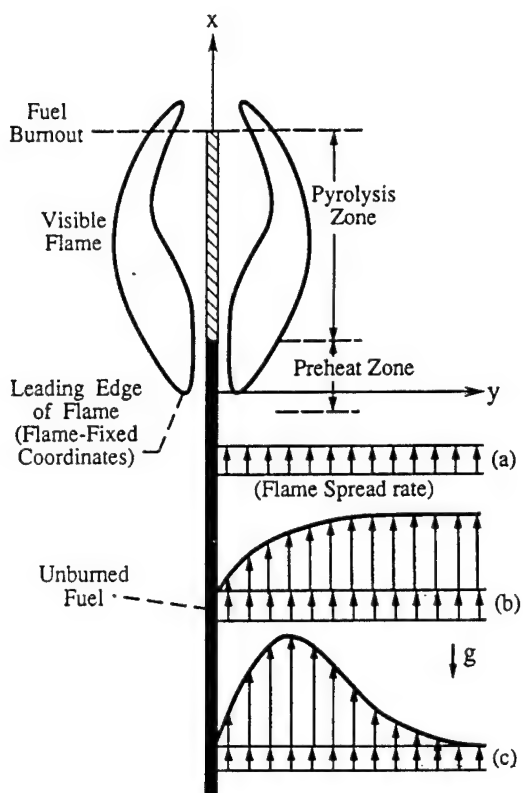


FIG. 26. Schematic of flame propagation with different opposed flow velocity profiles in the flame-stationary reference frame: (a) μg stagnant environment, (b) μg forced-convection environment, and (c) stagnant buoyant environment.

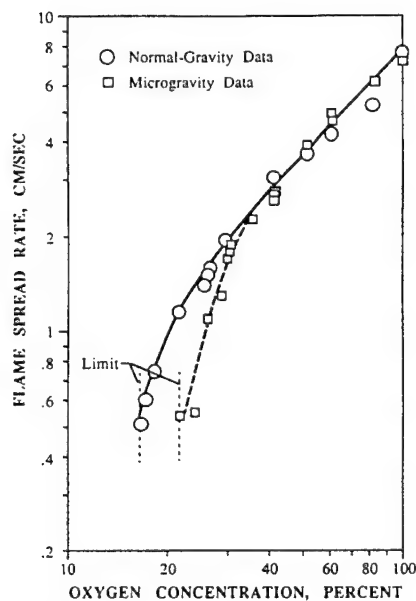


FIG. 27. Flame spread rates at ng and μg as a function of ambient oxygen concentration for a thin solid fuel. From Olson.¹⁵²

Recently, however, work has been initiated on flame spread over thick fuels,^{147,148} concurrent spread,^{147,149} and turbulent flames.¹⁵⁰ There is a growing body of literature on flame spread related to fire research (see Refs 146-163, and references cited therein, for additional information).

Figure 26 is a more detailed schematic showing an opposed spread configuration. Recognizing that the flame has a finite spreading speed, Fig. 26 shows that, in the flame-stationary reference frame, the flame experiences different upstream velocities against its motion if it is situated in (i) a zero-gravity, stagnant environment, (ii) a zero-gravity, forced convective environment, and (iii) a stagnant, buoyant environment. Spreading is accomplished through highly coupled convective-diffusive heat and mass transport, as well as radiative heat transport, near the leading segment of the flame. Here the forward diffusive and radiative heat transfer from the flame heats and pyrolyzes the solid fuel ahead and in the vicinity of the leading edge of the flame. The pyrolyzed fuel vapor then mixes with the fresh oxidizing gas from the freestream, forming a combustible mixture which subsequently reacts. Since the mixture around the leading edge of the flame is premixed, this flame segment resembles a premixed flame. Thus, spreading of such a global diffusion flame is achieved through propagation of this premixed flame segment that forms the leading portion of the flame. Fundamentally, we also recognize the similarity of flame spread to the stabilization of jet diffusion flames by a premixed flame segment. A comprehensive theory, yet to be formulated, should be able to describe both phenomena.

Microgravity experiments have been conducted in order to study flame spread without the complications from heat and mass transport processes associated with the induced buoyant flow.¹⁵¹⁻¹⁵⁵ It is also of particular interest to assess the influence of the convective environment on fire safety, especially under limit conditions. In general, it is found that as the external convective flow is reduced to values near zero, the flame temperature is reduced, and the flame becomes thicker and is located farther away from the surface. Figure 27 is an illustration of measurements of the flame spread rate over a thin paper sample as a function of the ambient oxygen concentration at ng and μg for a stagnant environment.¹⁵² It is seen that while the flame spread rate is basically not affected by buoyancy in highly-enriched oxygen environments, as the oxygen concentration is reduced the flame spreads more slowly at μg . Consequently, the flame fails to spread at a comparatively higher oxygen concentration at μg . The substantial difference between the spread rate at μg and ng also indicates the need to conduct these experiments at μg , as well as other intermediate gravity levels, in order to accurately assess the influence of buoyancy.

Figure 28 is an illustration of the flame spread velocity over thin paper samples as a function of the

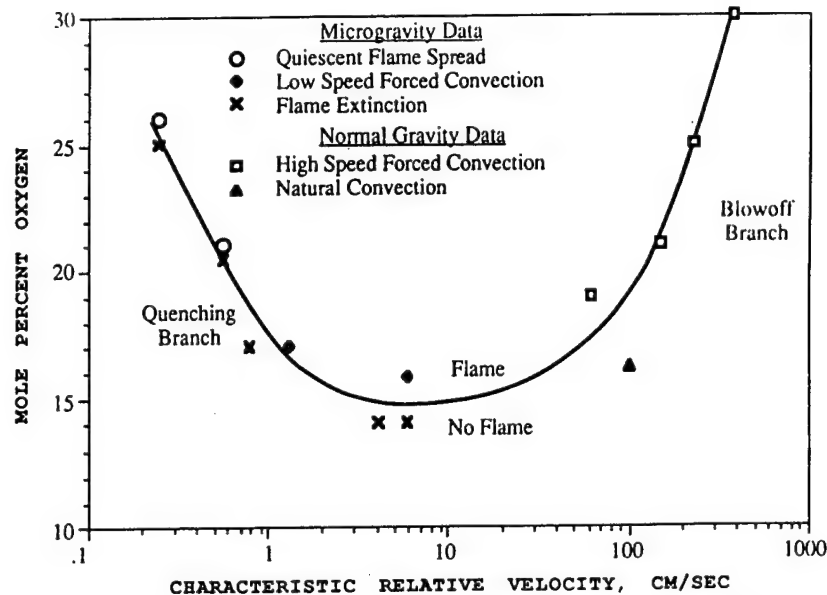


FIG. 29. Extinction boundary for flame spread over a thin fuel. From Olson.¹⁵²

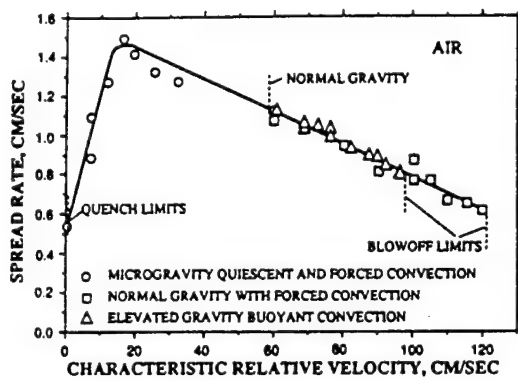


FIG. 28. Effect of opposed flow on the flame spread rate for a thin fuel. From Olson.¹⁵²

characteristic relative velocity experienced by the flame.¹⁵² The correlation considers all possible modes of convection, including quiescent and forced convection at μg , with and without forced convection at ng , and buoyant convection at elevated gravity. It is seen that the spreading response is nonmonotonic, with the spread rate exhibiting a peak value at a certain characteristic relative velocity. Such a nonmonotonic behavior is further demonstrated in Fig. 29, which is a plot of the limit of flame spread in terms of the oxygen concentration and the characteristic relative velocity. The results demonstrate that spreading is inhibited beyond a low and a high limit of the relative velocity. It is suggested¹⁵² that in the high velocity limit, relevant to buoyant or highly forced convective situations, the flame is simply blown off because a balance cannot be maintained between the local flow velocity and flame propagation speed at the leading edge of the flame. In the low velocity limit, relevant to stagnant μg situations, the reduced

convective rate of oxygen supply causes the burning rates, and thereby the heat release rates, to be too slow relative to that of the radiative heat flow, and the flame is quenched. The possible existence of such a dual extinction behavior was originally suggested by T'ien and Foutch and T'ien,¹⁵⁶⁻¹⁵⁸ based on numerical solutions of counterflow diffusion flames. Recent analytical studies of nonpremixed and premixed flames with flame and surface radiation^{159,160} also substantiate such a phenomenon.

The above results demonstrate that the complete suppression of convective flows does not necessarily lead to the most favorable situation to retard flame spread. In fact, it is seen that flame spread can be significantly promoted in a μg environment having a gentle breeze, somewhat lower than the lowest possible characteristic velocity for opposed-flow flame spread at ng , which is typical of the working environment in spacecraft. It is, however, also shown that an absolute minimum oxygen concentration of about 15% exists, below which opposed-flow flame spreading is not possible, at least for the materials used in the experiments of Fig. 29. These results are highly significant for fire prevention considerations and will be discussed again in Section 7.2.

The opposed-flow spreading of flames over thin fuels has also been numerically simulated.^{154,161-163} The predictions appear to quantitatively agree well with measurements at μg , especially at the leading edge.¹⁵⁴ Such agreement emphasizes the importance of the leading edge region and radiative transfer on opposed-flow flame spread.

A schematic diagram illustrating flame spread over a pool of liquid fuel appears in Fig. 30. The reference frame in Fig. 30 is the flame, so that both air and liquid approach the flamefront, with local modifications of surface tension and effects of buoyancy

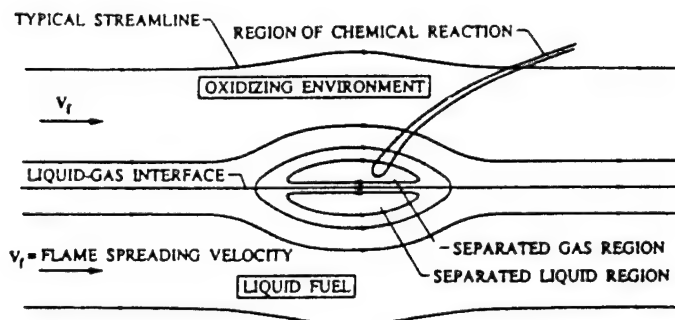


FIG. 30. Schematic showing flame spread over a liquid surface in the flame-stationary reference frame, with spreading being aided by surface tension. From Sirignano.¹⁶⁴

contributing to the formation of recirculation or separated-flow zones in both the gas and liquid. In this reference frame, the region near the leading edge of the flame resembles the flame attachment point of gaseous jet diffusion flames, discussed earlier. Thus, the leading edge involves a small premixed region fed by vapor leaving the fuel surface in its vicinity, in the same manner as that of flame spread over a solid surface discussed previously.

Flame spreading over the surface of a pool of liquid fuel (Fig. 30) possesses several distinctive characteristics which are different from the behavior of flames for solid fuels. First of all, since liquid fuels are more volatile than solid fuels, substantial amounts of fuel vapor can be generated in the region near the leading edge of the flame which controls the flame spread rate. Flame propagation is therefore sensitive to the vapor pressure (and thus the temperature) of the liquid, leading to the use of the flash point of the liquid fuel as an indicator of the different modes of spreading. It is reasonable to expect that with a moderate increase of the fuel vapor concentration, the premixture at the leading edge will be rendered more combustible and hence will propagate a flame faster when compared with a solid fuel. However, excessive enrichment could make the leading edge too fuel rich and consequently retard the spreading rate. Such a dependence has not been sufficiently studied.

There are two main reasons for the study of flame spreading over liquid pools at μg . First, since the surface tension in the flame region is now much smaller than that far upstream of the flame, because of the higher liquid temperature in the flame region, thermocapillary forces induce a surface velocity in the direction of flame propagation.¹⁶⁴ This effect subsequently increases the overall flame spread rate. Since the inherent flame spread rate in a stagnant μg environment is very small, as just shown, this surface tension-induced flow can become the controlling factor in flame spreading. The second consideration is that if the pool is sufficiently deep, then buoyancy will also affect the fluid motion within the liquid, and thus the flame spread rate. A complete understanding of the flame propagation process will

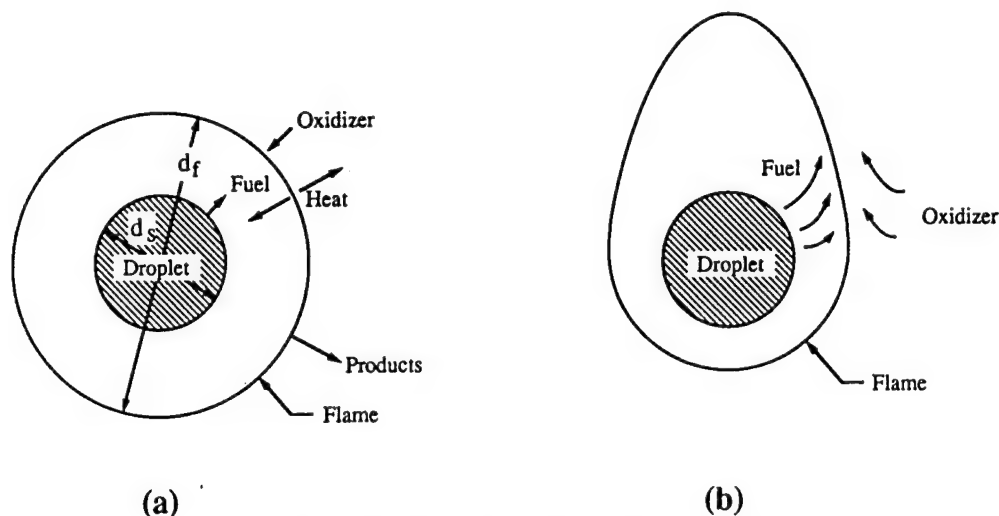
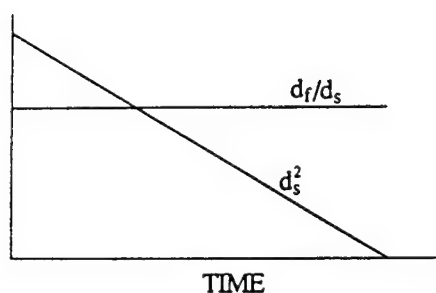
therefore involve studies of both gas- and liquid-phase processes.

Microgravity experiments using both shallow and moderately deep pools^{165,166} have shown that when flames at ng spread uniformly, the spreading rates at μg are almost identical to those at ng . This implies a minimal influence of buoyancy on the motion in either liquid or gas phases for this horizontally-oriented fuel surface. The spreading behavior, however, is markedly different under limit conditions of, say, low liquid temperatures. For these situations, spreading assumes a pulsating mode at ng , and is not possible at μg . It is reasoned that at ng a gas-phase recirculation cell is present ahead of the leading edge, being formed by the concurrent gas flow next to the liquid surface due to the no-slip boundary conditions and the opposed flow above it induced by buoyancy.¹⁶⁷ The gas-phase recirculation cell also induces a liquid-phase recirculation cell. The gas-phase recirculation cell facilitates fuel-air mixing, especially near limit conditions, and therefore promotes combustion. At μg , such a recirculation cell may not exist (due to hot gas expansion) and flame spreading is therefore not possible under the same limit conditions. However, when an opposed flow is externally applied at μg , the flame is sustained under otherwise extinguished conditions.¹⁶⁶

6.3. Droplets

Droplet vaporization and combustion is an essential process during spray combustion in liquid-fueled engines and combustors. It is also a classical model problem of heterogeneous combustion because, by assuming quasisteadiness and spherical symmetry, the mathematical aspects of the problem can be greatly simplified.

The basic droplet combustion model,¹⁶⁸ depicted in Fig. 31, describes the gasification of the liquid fuel at the droplet surface and its subsequent outward transport to meet inwardly-diffusing oxidizing gas in a thin flame region. The classical d^2 law, derived by assuming spherical symmetry (Fig. 31a), quasisteadiness, and flame-sheet combustion, then predicts that

FIG. 31. Schematic of drop envelope flames at (a) μg , and (b) ng .FIG. 32. Theoretical predictions of spherically symmetric drop combustion according to the d^2 law.

the square of the droplet diameter, d_s^2 , decreases linearly with time, and that the ratio of the flame diameter, d_f , to the droplet diameter, d_f/d_s , is a constant, as illustrated in Fig. 32.

Numerous attempts have been made to verify the d^2 law. Normal gravity experimental results show that d_s^2 indeed varies approximately linearly with time, with the deviation from linearity depending on the extent of second-order influences. Attempts to quantify the behavior of the flame size, however, proved futile because it is meaningless to define a flame diameter when the flame is severely elongated by the presence of buoyant flow (Fig. 31b). Furthermore, since the intensity of buoyancy continuously changes because of the steadily decreasing droplet size, an unsteady effect is also introduced. Thus, μg experiments are needed to better understand droplet combustion.

Kumagai and coworkers^{169,170} pioneered the use of the drop tower to study droplet combustion at μg . A spherically-symmetric droplet combustion configuration was observed during these experiments. The period of free-fall in Kumagai's experiments, however, was fairly short and therefore only covered the early portion of the droplet lifetime.

An alternative approach to minimize buoyancy during droplet burning is to conduct the experiment in ng environments at reduced pressure,⁸ down to about 0.1 atm., as mentioned earlier. Since droplet burning is a diffusion-controlled process, and since the density-weighted mass diffusivities are pressure insensitive, this approach is suitable to study certain small-scale (say 1 mm or less), low-velocity, steadily-burning diffusion flame phenomena. The reduced chemical reactivity can also be partially compensated by using oxygen-enriched environments, which have the additional advantage of reducing the flame size and hence the extent of buoyancy experienced by the flame. Results from such a study show that while d_s^2 still varies approximately linearly with time, the behavior of the flame diameter completely disagrees with the d^2 law prediction.⁸ Specifically, the flame-front standoff ratio d_f/d_s continuously increases with time for low to moderate ambient oxygen concentrations, but increases and levels off for higher ambient oxygen concentrations (Fig. 33). This behavior has recently been substantiated by experiments conducted at μg .^{171,172} Theoretical results⁸ showed that this behavior is caused by transient phenomena, that have been called "fuel vapor accumulation effects." That is, at the moment of ignition, the amount of fuel vapor in the region interior to the flame is low compared with the d^2 law value, being constrained by the amount which existed for the vaporizing droplet prior to ignition. Thus, subsequent to ignition, only part of the fuel that is gasified is used to support burning at the flame, the rest is used to build up the fuel vapor concentration in the inner region of the flame. The flame diameter consequently grows with increasing amounts of fuel vapor accumulation. The growth of d_f/d_s continues with time for low ambient oxygen levels where the flame is relatively large, while it levels off for higher ambient oxygen levels where the flame is relatively small and

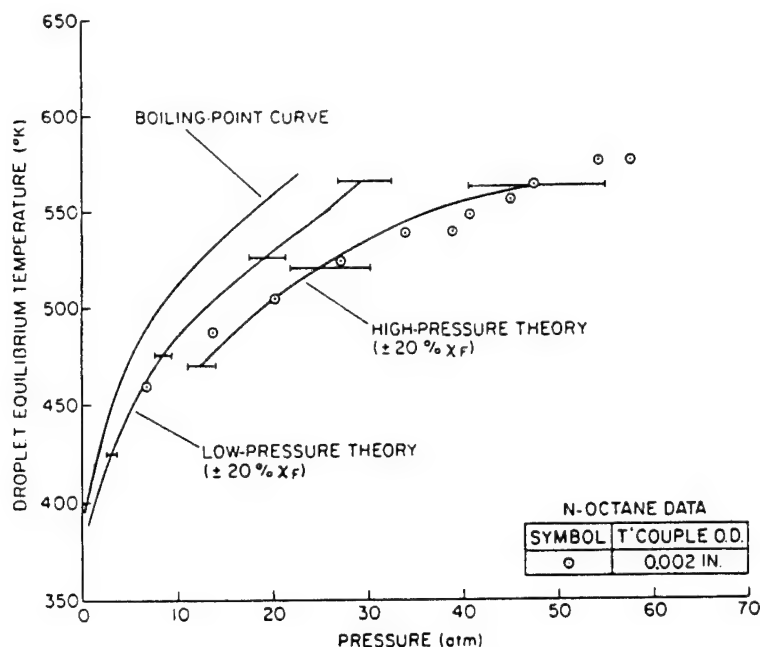


FIG. 34. Theoretical and experimental steady-burning temperatures of *n*-octane drops burning in air at low gravity conditions. From Lazar and Faeth.¹⁷⁴

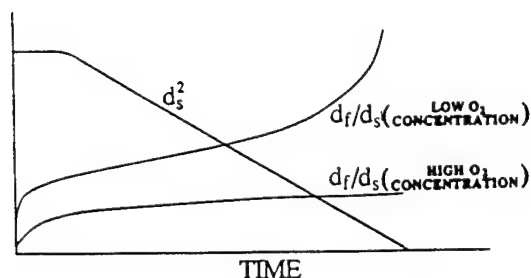


FIG. 33. Behavior of spherically symmetric drop combustion demonstrating the effect of fuel vapor accumulation on flame sheet location.

there is less need for fuel vapor accumulation. The phenomenon can be completely described by considering gas-phase quasisteadiness, treating the fuel vapor accumulation process at the same slow timescale as that of droplet surface regression. Indeed, by suppressing fuel vapor accumulation, through the use of the d^2 law concentration profile as the initial condition, a separate analysis¹⁷³ allowing for gas-phase transient diffusion shows that fuel vapor accumulation has only minor influence on the flame response. Further discussion of this effect can be found in Ref. 168.

Gas-phase unsteadiness is also important during droplet vaporization and combustion in high-pressure environments when effects of forced convection are absent, because the droplet approaches its thermodynamic critical point where the liquid and gas densities are comparable. When the liquid reaches or exceeds its critical point, liquid properties are lost,

the droplet acts like a puff of gas, and it is meaningless to talk of droplet combustion. Experiments at low gravity have been used to identify droplet properties at high pressures and the conditions for the onset of supercritical combustion.¹⁷⁴ These experiments involved igniting droplets within a pressure vessel at various ambient oxygen concentrations and using a free-fall facility to provide low-gravity conditions. It was found that the increased solubility of the liquid to gases at its surface at high pressures has an important effect on droplet surface properties and the conditions required for supercritical combustion. Some typical results are illustrated in Fig. 34, which is a plot of the liquid temperatures during the steady burning phase (where the droplet has reached its wet bulb temperature) as a function of the ambient pressure for *n*-octane droplets burning in air. The plot includes measured liquid temperatures, the standard boiling point curve and predictions both ignoring (low-pressure theory) and considering (high-pressure theory) effects of dissolved gases in the liquid. The plots are terminated at high pressures at the point where supercritical combustion conditions are reached. The high-pressure theory is in reasonably good agreement with the measurements, and shows that the dissolved gases tend to reduce liquid temperatures at each pressure. Additionally, the dissolved gases also tend to increase the ambient pressure required for the droplet to exceed its thermodynamic critical point, beyond both the critical pressure of the pure fuel and the predictions of the low-pressure theory.¹⁷⁴

Effects of unsteady combustion in convection-free environments at high pressures have also been stud-

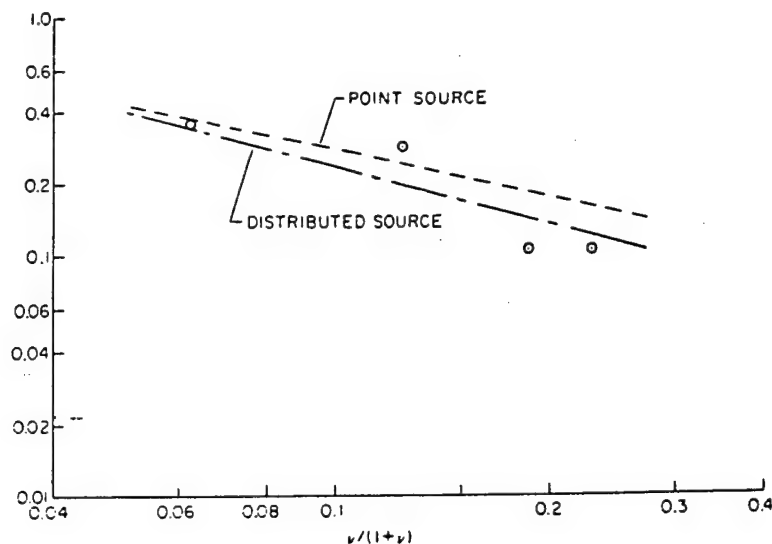


FIG. 35. Supercritical combustion lifetimes of *n*-decane at 68 atm, as a function of stoichiometric parameter (corrected to an initial drop diameter of 0.875 mm). From Faeth *et al.*¹⁷⁵

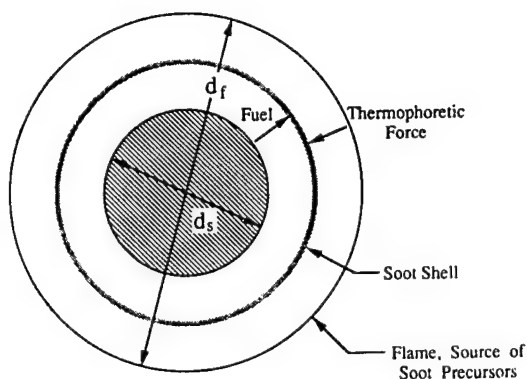


FIG. 36. Schematic showing the soot formation phenomenon during combustion at μg .

ied at low-gravity conditions, using a pressure vessel in a free-fall facility.¹⁷⁵ Combustion lifetimes (defined as the time between the droplet exceeding its thermodynamic critical condition and the end of combustion) were measured at high pressures where most of the combustion process occurs after the droplet exceeds its thermodynamic critical point. These measurements were compared with predictions of Spalding¹⁷⁶ and Rosner,¹⁷⁷ who treated unsteady combustion from point and distributed sources of fuel vapor, respectively. Predicted and measured results for *n*-decane droplets at 68 atm. and various ambient oxygen concentrations are illustrated in Fig. 35, where $v = Y_{O_2}/\sigma$, and σ is the stoichiometric mixture ratio. The distributed source theory yields better agreement with the measurements, but there is little to choose between the two in view of uncertainties in estimating transport properties in flame environments. Additionally, predictions of the maximum

flame radius during combustion were in good agreement with measurements for various ambient oxygen concentrations and pressures.¹⁷⁵ Subsequently, additional experiments on high-pressure droplet combustion at μg have been completed, as well as the development of advanced theories allowing for unsteady and real-gas effects in both cases (see Shuen *et al.*¹⁷⁸ and Faeth¹⁷⁹ for discussion of this work). These reviews indicate that much remains to be done to resolve the thermochemical and transport issues of droplet vaporization and combustion at the near-critical and supercritical conditions that are relevant to numerous high-pressure combustion processes. Thus, significant opportunities remain to exploit the demonstrated capabilities of combustion at μg for studies of high-pressure droplet combustion phenomena.

Recent μg experiments¹⁸⁰⁻¹⁸² have also revealed an unexpected, interesting aspect of droplet combustion, namely the formation of a soot layer between the droplet and the flame (Fig. 36). This phenomenon is caused by the back-diffusion of soot particles that are formed near the flame toward the droplet as a result of thermophoresis. Since this back-diffusion is opposed by the outwardly-transported fuel vapor, the soot particles eventually stagnate before reaching the droplet surface. The continuous accumulation of these soot particles eventually forms the shell as observed. The role of gravity here is apparent: soot layer formation, as seen at μg , has not been observed at ng for envelope flames, with such 'clean-burning' hydrocarbons as *n*-heptane in the presence of even very weak convective motion. Apparently the weak convection continuously sweeps the soot precursors and particles towards the flame segment in the wake on the leeward side of the droplet, causing their burnout.¹⁸³ This phenomenon can have a profound

influence on our understanding of soot formation with and without external convection. It is also of interest to reassess the sooting tendencies of those fuels which are usually considered to be nonsooty.

Another interesting phenomenon observed during recent μg single-component droplet combustion is that the droplet burning rate appears to be substantially lower than previously observed. Specifically, the classical result of Kumagai *et al.*¹⁷⁰ showed that the burning rate constant of *n*-heptane droplets in air is $0.78 \text{ mm}^2/\text{s}$, while the recent determination¹⁸¹ showed a lower value of around $0.6 \text{ mm}^2/\text{s}$, which also decreases with increasing initial droplet size.¹⁸⁴ The tentative explanation for this burning rate retardation is that soot formation acts as a chemical heat sink in reducing the total amount of fuel mass to be oxidized, and that the soot shell could also act as a partial barrier for heat transfer from the flame to the droplet surface. The fact that Kumagai *et al.*¹⁷⁰ obtained a higher burning rate, and indeed did not observe any soot shell at all in their μg experiments, was postulated to be caused by the dispersion of the soot particles by the convective motion formed when the suspension fiber was abruptly pulled at the start of their particular experiment. Results of this nature demonstrate the sensitivity of μg experiments to small currents of air movement and the care needed to prevent their occurrence.

Spherically-symmetric droplet combustion also serves as a good model problem for the study of diffusion flame reaction kinetics.¹⁸⁵⁻¹⁸⁸ Recently there has been considerable effort to extract fundamental kinetic information by comparisons between theoretical and experimental results for certain flame processes. The accuracy of these comparisons, however, is frequently compromised by inaccurate descriptions of the flow field. This difficulty does not exist for the simple spherically-symmetric droplet combustion configuration. Since droplet burning is basically diffusion controlled, kinetic effects are best manifested by the ignition and extinction events. In particular, recent studies at μg have made frequent observations of drop extinction before all the liquid has evaporated.^{172,182,187,188}

A complication that can result from droplet studies of flame kinetics is the potential influence of liquid-phase processes on gas-phase behavior. For example, when methanol is used as the fuel, it can potentially dissolve the water generated at the flame.^{190,191} Thus, shortly after ignition, the droplet becomes a bicomponent mixture, whose amount and spatial distribution are not only unknown but also vary with time. While these quantities could conceivably be determined by numerical simulation, it is difficult to be quantitatively accurate due to the uncertainties of liquid-phase transport properties. Due to the complications as a result of water dissolution, the reliability of determining the gas-phase kinetics is correspondingly compromised.

A droplet ignition study at μg has also been con-

ducted by rapidly submerging a drop in a heated gas within a furnace, with the apparatus contained in a free-fall facility, yielding ignition delay times for various liquid fuels and ambient conditions.¹⁹² This test configuration provides a convection-free environment that simplifies both kinetic analysis and the interpretation of measurements which deserves additional consideration.

Several μg multicomponent droplet combustion programs are also currently underway. The most important factors which distinguish multicomponent droplet combustion from pure liquid droplet combustion are that the volatility differentials among the various components are an essential factor, and that liquid-phase mass diffusion can be very slow compared with either the surface regression rate or liquid-phase thermal conduction.^{168,193,194} The parameter which characterizes the efficiency of liquid-phase mass diffusion is a Peclet number, Pe , defined as the ratio of the surface regression rate to the liquid mass diffusivity. For droplet combustion, Pe is usually a large number because liquid-phase mass diffusivities are small. Consequently, after the initiation of gasification of an originally uniform droplet consisting of, say, two components having different volatilities, there exists an initial period when the volatile component in the surface layer is preferentially gasified and the droplet temperature is controlled by the boiling point of this volatile component. This preferential gasification, however, will eventually slow down due to the reduction of the volatile concentration in the surface layer and the extremely slow rate with which it can be replenished from the core of the droplet. A steady state will then be reached at which the rate of supply of the components is equal to the rates with which they are vaporized. Furthermore, since the surface is now more concentrated with the less volatile, higher-boiling-point component, the droplet temperature will also become correspondingly higher. It is therefore clear that between the initial and the steady states a transition droplet heating period must exist, during which the droplet temperature increases while the droplet gasification rate becomes slower because most of the heat it receives is now diverted to effect droplet heating.

Experimental results at ng , but for weakly-convective situations, substantiate this three-stage behavior.^{195,196} Figures 37 and 38 respectively show the variations of d_t^2 and the normalized flame diameter with time for a bicomponent droplet initially consisting of 70%-heptane and 30%-hexadecane. The existence of the transition heating period, characterized by a slow gasification rate and a simultaneous shrinking of the flame diameter, is particularly noteworthy. The above behavior has also been observed during recent microgravity experiments.¹⁹⁷

The combustion of multicomponent droplets is sometimes accompanied by a microexplosion phenomenon, which involves either mild fragmentation

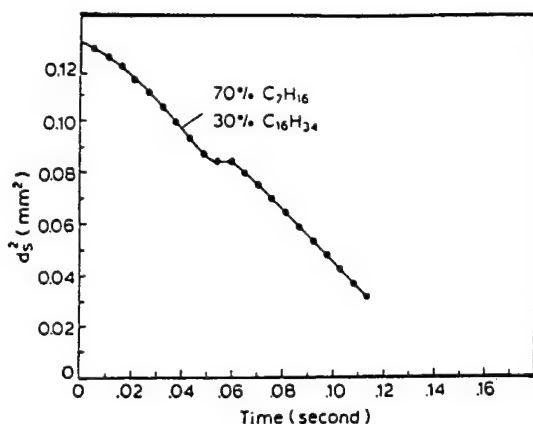


FIG. 37. Temporal variation of the droplet diameter for heptane-hexadecane droplets burning in air. From Wang *et al.*¹⁹⁵

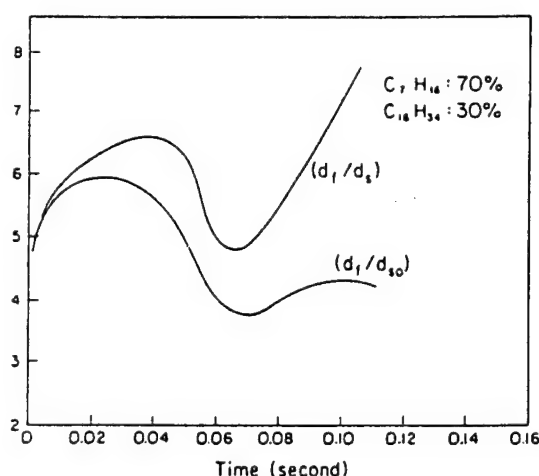


FIG. 38. Temporal variations of normalized flame-diameter (d_f/d_{s0}) and the flame-front standoff ratio (d_f/d_s) for a 70% heptane-30% hexadecane droplet burning in air. From Wang *et al.*¹⁹⁵

or violent rupturing of the droplet as it burns. The fundamental cause of microexplosion is the superheating of the more volatile components trapped in the inner core of the droplet whose temperature is controlled by the higher boiling points of the less volatile components. Thus, when these more volatile components are locally heated to the limit of superheat, instantaneous gasification and thereby droplet rupturing occurs. The tendency to microexplode has been experimentally observed at ng, but with weakly-convective environments,¹⁹⁸ to increase with increasing system pressure, due to the corresponding increase of the liquid boiling point and the nonideality of the mixture. These results were again recently confirmed by μg experiments.¹⁹⁹

A heterogeneous burning problem which is somewhat similar to droplet combustion is the candle flame,^{200,201} in which a diffusion flame is established between the inwardly-diffusing oxidizer gas and the

outwardly-diffusing vaporized wax. Since the length of the wick is fixed, the phenomenon is basically steady, in contrast with quasisteady droplet burning situations in which the droplet size continuously shrinks. At μg , the flame appears to be somewhat spherical, in spite of the axisymmetric nature of the fuel source, probably due to the large flame size in comparison with the length of the wick. However, the soot shell which was present for the droplet flame was not observed for the μg candle flame. More experiments are needed to draw conclusions about the characteristics of these flames.

The combustion of metals at μg has also received some attention, motivated by concerns about hazards associated with high-pressure oxygen systems in space applications.²⁰² This work involved the combustion of rods of aluminium alloy, stainless steel alloy, iron and titanium, as well as sheets and meshes of stainless steel alloy. Combustion was studied in pure oxygen at various pressures. The experiments involved upward propagation at ng, in order to avoid irreproducible results due to molten metal flowing along the sample surface, as well as tests at μg . The combustion of these materials involves the formation of molten balls at the end of the sample, which serve as the fuel source for the flame. It was found that the regression rate of the melting interface was significantly greater at μg than ng and that some sample shapes, such as thin sheets, that extinguish at ng burn to completion at μg . Much remains to be done, however, in order to exploit the use of μg to study metal combustion: in particular, the high melting and boiling points of metals and their combustion products, their capabilities to form complex condensed phase mixtures, and the variety of combustion mechanisms for various metal-oxidant combinations, provides a host of theoretical and experimental problems.

The combustion of solid materials other than metals has received only limited attention at microgravity, for example, the study of the combustion of carbon rods by Spunkler,²⁰³ and additional work should be encouraged. In particular, investigation of solid particle combustion, analogous to drop combustion, should provide valuable background information needed to gain an understanding of more complex processes such as flame spread along surfaces and heterogeneous flame propagation through suspensions.

7. SPACECRAFT FIRE SAFETY

7.1. Introduction

The previous discussion has amply demonstrated that the flame spread rates and flammability limits at μg and ng are not the same, and that information gained during ng studies may not be accurately transferable to μg situations even as limiting case

criteria. For example, no solid or liquid material has yet been studied to find the lean flammability limits (or the limiting oxygen concentration for flame propagation) in all the obvious configurations, for example, upward and downward flame spread at ng, and quiescent, opposed and concurrent flow at μg ,²⁰⁴ therefore, it has not been possible to quantitatively demonstrate the relationship between fire safety properties at ng and μg . Furthermore, μg flame sizes are expected to be large while the environment on board manned space facilities such as the space station is compact, both of which tend to aggravate the consequence of a fire. Thus, specific information relevant to spacecraft fire safety is discussed in the following, treating flame-inhibiting atmospheres, fire detection and fire extinguishment, in turn.

7.2. Flame-Inhibiting Atmospheres

The control of fires within isolated inhabited compartments such as spacecraft, aircraft cabins, submarines, medical chambers, etc. involves novel problems in comparison with conventional systems.²⁰⁵⁻²¹⁸ In particular, any fire must be prevented or extinguished in a way that does not endanger the occupants, interfere with operations, or damage operating systems in a very confined environment. This is particularly critical for spacecraft operations because the occupants may have to live and function in the postfire environment for a considerable period of time. Some of the problems of fire control for these circumstances will be considered subsequently; the following discussion will address available tactics for the prevention of fires, emphasizing the potential use of flame-inhibiting atmospheres that might be able to entirely eliminate fire hazards in spacecraft.

The most common method of fire control is to eliminate potential ignition sources and to select materials that minimize the risk of incidence of fires, for example, 'design to preclude' fire.^{207,211} This is effective up to a point, however, the potential for fires cannot be completely eliminated in this manner and the required modifications of system design and materials are costly and inevitably involve excessive compromises of performance and flexibility of spacecraft operations.^{205,207} These problems will only increase as the exploitation of space proceeds and a greater range of manned spacecraft operations are undertaken. Thus, more effective alternatives must be sought.

The fact that the environment within spacecraft is completely under human control offers a novel approach toward avoiding fires by using flame-inhibiting, or 'fire-safe,' atmospheres as an approach toward design to preclude. In particular, a property of fire-related phenomena is that they tend to be functions of the *fractional* amount of oxygen in the

atmosphere. In contrast, human comfort and performance seem to depend mainly on the *absolute* amount of oxygen in the atmosphere (or the oxygen partial pressure). Thus, the potential exists to find a composition of the atmosphere that will not support combustion but that will support normal human activities indefinitely. This approach has been advocated by Carhart,²⁰⁸ based on experience with undersea systems such as submarines and Sealab.³

Underwater systems provide several examples of fire-safe atmospheres.^{207,208,210} For example, the atmosphere within Sealab was rendered completely fire safe by pressurizing the vehicle to 7 atm., while keeping the partial pressure of oxygen at the standard sea level value of 0.21 atm. This reduced the oxygen mole fraction to only 3%, which is well below flammability limits, both on Earth and in space. This was a very natural solution for Sealab because normal operations required the pressurized atmosphere in any event; however, this specific technique is not immediately transferrable to spacecraft, where similar pressure increases would cause substantial structural weight penalties. Nevertheless, it seems likely that similar fire-safe atmospheres, having operationally acceptable properties, can be found for spacecraft. They certainly should be sought due to the enormous advantages of eliminating fire hazards for manned space activities, with minimal limitations on materials properties.

Fire-safe atmospheres for spacecraft would be unique from other applications; nevertheless, existing information on flammability limits, studies of fire-safety parameters such as minimum oxygen indices, and specific studies of flame-inhibiting atmospheres, can provide some general guidelines.^{3,205-211} For example, the flame spread limit illustrated in Fig. 29 for thin fuels at μg is roughly 15% oxygen by volume. On the other hand, it is generally agreed that an oxygen partial pressure of 16.5 kPa, equivalent to an altitude of 1800 m, is acceptable for long-term human activities,²¹¹ for example, such conditions are comparable with those of high-altitude cities such as Mexico City and Denver. Thus, satisfying both requirements with an oxygen-nitrogen atmosphere would imply a total pressure of 118 kPa. Another reduced oxygen atmosphere was identified by studies sponsored by the Navy that showed acceptable performance of spacecraft crews in an atmosphere having an oxygen partial pressure and mole fraction of 16.0 kPa and 11%.²¹² This mole fraction is thought to retard most fires, based on ng data, and would require a total pressure of 145 kPa. These reduced-oxygen atmospheres are not major departures from the atmosphere planned for space stations; namely, air at atmospheric pressure (101.3 kPa).²¹³ Naturally, these prescriptions are only suggestive because limited data for a single material and ignition mode, etc. such as the results of Fig. 29, or test results limited to ng, are certainly not sufficient to make a judgement concerning fire-safe ambient oxygen mole fraction levels at

TABLE 2. Fire-safe atmosphere tests at atmospheric pressure^a

O ₂ (% by vol)	Inert (% by vol)	C _p (kJ/kg O ₂ K)	Fuel	Remarks
40.5	59.5 C ₂ F ₆	5.77	filter paper	ignition flame only
21.1	20.2 C ₂ F ₆ , 58.9 He	5.91	filter paper	ignition flame only
			raw cotton	
			polyurethane foam	
21.0	10.0 C ₄ F ₆ , 69.0 He	5.68	filter paper	ignition flame only
21.0 ^b	21.0 C ₂ F ₆ , 69.0 N ₂	6.68	filter paper	ignition flame only
			cotton cloth	
			polyurethane foam	
			kerosene	
16.9	19.0 CF ₄ in air	6.49	cotton flannel	no combustion
18.4	12.0 C ₂ F ₆ in air	6.49	cotton flannel	no combustion
19.0	9.0 C ₃ F ₈ in air	6.51	cotton flannel	no combustion
17.8	15.0 CF ₄ in air	5.95	kerosene	no combustion
19.2	8.0 C ₂ F ₆ in air	5.72	kerosene	no combustion
19.4	7.0 C ₃ F ₈ in air	5.99	kerosene	no combustion

^aFrom Huggett.²⁰⁵^bElectrically heated solid rocket propellant as ignition source; remainder ignited using an electrically-fired match.

μg. Additionally, there are numerous human factors, and spacecraft design and operational requirements, that must be taken into account. Nevertheless, these very preliminary results do suggest that potentially fire-safe atmospheres may not be particularly unusual or hard to provide.

Research carried out by Huggett^{205,206} provides other general guidelines for selecting fire-safe atmospheres. In particular, he found that flame spread over solid surfaces could be correlated with the heat capacity of the gaseous environment per mole of oxygen, and that when this heat capacity reached the range 5–6 kJ/kg O₂ K (40–50 cal/g mol O₂ K) organic fuels ceased to burn at atmospheric pressure. This prompted the study of fully fluorinated compounds of carbon as diluents for suppressing combustion because they have unusually high heat capacities among known inert gases.²⁰⁵ Table 2 gives a summary of data on atmospheres that did not support combustion for various fuel samples ignited by either an electrically fired match, or a small piece of solid propellant, placed at the bottom of the sample at atmospheric pressure. All of these mixtures are characterized by a specific heat capacity in the range 5.7–6.7 kJ/kg O₂ K, while mixtures having lower specific heats sustained combustion. The results also indicate flexibility for meeting the specific heat criterion: some mixtures involved O₂ concentrations greater than normal atmospheric levels, while the use of helium as a diluent yielded overall molecular weights and densities similar to normal air. Additionally, limited testing with animals—mice, rats, rabbits and dogs—did not indicate significant physiological problems when subjected to at least some of these environments.²⁰⁵ Other saturated fluorine compounds, such as sulfur hexafluoride, offer similar advantages.²⁰⁷ A potential problem with fully fluorinated compounds, similar to those listed in Table 2, however, involves potentially harmful emissions of HF in the confined fire environments of spacecraft.²¹⁵

The previous results suggest that there is potential for fire-safe atmospheres with acceptable properties for spacecraft operations. However, available information is woefully inadequate in view of the importance of this selection. Obvious test variables such as fuel type, fuel configuration, method of ignition, ignition energy, flow properties, the presence of a μg environment, and the effect of total pressure have not been examined. Furthermore, fundamental understanding of flammability is incomplete so that extrapolation of limited data is unreliable, as discussed earlier. Finally, effects of fire-safe atmospheres on the performance and health of humans and other biological systems, on the quality of voice communications due to changed acoustic properties, and on spacecraft design and operations, certainly have not been adequately evaluated. In view of the potential importance of fire-safe atmospheres to manned space flight, however, this problem clearly merits a comprehensive interdisciplinary research program involving specialists in combustion, space medicine, human performance, and spacecraft design and operations, in order to resolve these issues. This task represents long-term research and it is doubtful that the results could apply to current plans for Space Station Freedom where the decision to use air at atmospheric pressure was made some time ago, based on a desire to have an atmosphere similar to sea level air so that ground data can be compared directly with μg flight data (see J. H. Kimzey in Appendix C of Ref. 218). Nevertheless, it would be desirable to gain a better understanding of fire-safe atmospheres for application to manned space activities in the future.

7.3. Fire Detection and Extinguishment

Problems of fire detection and extinguishment in spacecraft have been recognized for some time,

prompting several recent studies and reviews of this technology (see Refs 207 and 214 and references cited therein). Methods of fire detection have progressed from the crew serving as fire detectors during the first manned space programs, to multiple ionization smoke detectors in the Space Shuttle, to proposals for multiple smoke, UV flame and thermal detectors for Space Station Freedom.²⁰⁷ The main concerns are the slow rate of response of detectors at μg (except for radiation sensors), the modified convection patterns of flames at μg as opposed to ng , and the requirements for high sensitivity to compensate for the relatively slow response of μg environments to smoldering and flames. Locating detectors to exploit the spacecraft ventilation system helps to mitigate these difficulties to some extent, but fault conditions where the ventilation system is inoperative, must still be addressed. Finally, the technology base for fire detection in spacecraft is very limited due to the difficulties and costs of arranging full-scale tests at μg .

Problems of fire extinguishment in the confined environment of spacecraft were mentioned earlier in connection with fire-safe atmospheres. Thus, the use of techniques developed for underwater applications, by temporarily flooding the environment with an inert atmosphere, deserve consideration.^{207,208,210} For example, Halon 1301 effectively prevents flammability at concentrations less than 6%. Additionally, Navy applications, where fluorinated compounds are not acceptable due to problems with the life support system, have used nitrogen flooding for years.^{208,210} A better understanding of systems of this type for spacecraft at μg is closely related to research on fire-safe atmospheres, and merits study on the same basis.

Fire extinguishing systems on early manned spacecraft were primitive, involving the backup use of the water dispenser and aqueous gels.²⁰⁷ This has evolved to multiple Halon 1301 dispensers on the Space Shuttle, similar to aircraft fire protection systems.^{207,214} Unfortunately, Halon 1301 is less effective for smoldering fires (which are a common scenario for fires in spacecraft) and presents environmental problems due to emissions during ground-based activities (alternatives are being sought for aircraft as a result), as well as potentially harmful postfire emissions of HBr and HF.²¹⁵ The use of other gases, such as carbon dioxide which has been proposed for the space station, and venting the space where combustion is occurring, are being studied. Problems with less effective extinguishing agents, or venting, involve enhanced combustion, at least for a short time, due to increased convection from forced flows associated with the operation of these systems.²⁰⁷ Thus, much remains to be done in order to establish a proper technology base for fire extinguishment on board spacecraft, even for relatively near-term applications such as the Space Station.

8. MICROGRAVITY FACILITIES

8.1. Introduction

Experimental research under controlled gravity conditions is constrained in many ways. The most obvious constraints are the small number of facilities that can provide low gravity conditions, the long development times required to prepare experiments for these facilities, the large costs associated with the use of the most flexible facilities, the limitations concerning the instrumentation that can be used, the available test times at low gravity, and the availability of the facility to experimenters. These issues, as well as the capabilities of available facilities, will be discussed in the following sections, considering drop towers, aircraft facilities, sounding rockets and spacecraft, in turn.

8.2. Drop Towers

Drop towers provide the easiest access to a μg environment and many facilities of this type have been developed by individual workers. These facilities generally involve test times less than 1 s, which require free-fall distances less than 5 m. The capabilities of drop towers to sustain low-gravity conditions varies with the specific design, but it is not difficult to achieve values smaller than 10^{-3} g . Improved performance can be obtained by using a drag shield, where the actual test chamber falls within an outer freely falling chamber so that the relative velocities of the test chamber, and thus the drag forces, are minimized. A simpler approach, acceptable when only short test times are needed, is a short-drop tower where a drop shield is often not needed, for example, a 1.2 m free-fall distance that involves relatively small drag forces can provide 0.5 s test times at good low-gravity levels.

Longer test times require more sophisticated facilities with drop towers at NASA Lewis and elsewhere in the United States and Europe providing 2–5 s at μg down to 10^{-4} – 10^{-6} g , and a new facility in Japan providing 10 s test times at similar conditions (see Lekan²¹⁶ for a detailed description of major drop-tower facilities available in the United States). Additionally, these facilities can provide substantial test chamber volumes (10–1000 L). A disadvantage of most free-fall facilities having longer test times, however, is that the test apparatus is subject to a considerable shock load, of the order of 100 g, at the end of a drop test. The 10 s drop tower in Japan is an exception, however, and has relatively modest deceleration rates.

The facilities at NASA Lewis are representative of drop-tower technology for test times in excess of 2 s.^{5,216} Designs of test chambers, plumbing systems, data acquisition and control electronics, and film- and video-based imaging equipment are mature for

this environment, and a number of somewhat standard test rigs are being used in both 2.2 and 5 s drop towers. Quiescent chambers with volumes of ~ 40 L are available in which various atmospheric mixtures up to more than 2 atm. pressure can be formulated. These chambers have optical access principally designed for photography and video records and have electrical connections for conventional transducers. A combustion tunnel is being used in the 2.2 s drop tower. The tunnel provides an unvitiated blowdown flow of preformulated atmospheres at 1 atm. pressure, and flat inlet velocity profiles having velocities of 0–300 mm/s. The tunnel has been used for flame spreading experiments with solid fuels, but could be adapted for experiments in other combustion systems. The tunnel provides orthogonal views across the 200 mm diameter test section, which have been used for visualizing flames and cold forced flow. Development of a tunnel for the 5 s drop tower has been initiated.

Due to the rather severe impact loads, adapting common optical diagnostic methods to drop-tower experiments has been slow. Currently, workers at NASA Lewis are attempting to introduce a rainbow schlieren technique into a drop-tower package. Similar to conventional schlieren photography, refractive index gradients are detected, but are encoded here as variations in visible wavelength rather than the less-sensitive variation in monochromatic intensity. Rainbow schlieren methods will provide a visualization technique for the refractive index fields of μg flames. If constituent, and temperature influences on density can be distinguished, then the method could be used as a quantitative measure of temperature fields. Measurement of sooting characteristics of μg flames are being pursued in drop-tower experiments at NASA Lewis with the introduction of simultaneous absorption-scattering measurements and cold-probe thermophoretic sampling of soot particles. These measurements are intended to provide spatially distributed measurements of soot particle number density and size distributions.

Velocimetry at μg is being attempted in drop-tower experiments at NASA Lewis that will use particle imaging velocimetry (PIV) methods. In PIV, a sequence of computer enhanced images of thin, laser-illuminated sheets from within a flowfield are correlated to establish the path of seed particles and thereby their velocities. This method is limited in its ability to measure large velocities by the intensity of the illumination source and the sensitivity of the image detector. Current source and detector technologies were vulnerable to the impact loads experienced in the past in drop towers, which limited this measurement scheme to low-velocity flows (up to 100 mm/s) in, for example, liquid-fuel pools. Recently, however, drop rigs outfitted with shock isolation platforms have been developed, which reduce landing loads to 30 g, a level that a wide array of commercial instrumentation can sustain without damage. Thus, a

broader range of application of PIV, and other shock-sensitive instruments such as rainbow schlieren deflectometers, can be anticipated in the future.

Thus, at the current time, an interesting array of test facilities and instrumentation are available for μg combustion tests in drop towers, with more advanced laser diagnostics in the offing. A frustrating feature of these facilities, however, is that 2–10 s is a perilously short time to develop combustion processes and to achieve the steady-state conditions for combustion experiments that are easiest to interpret. This has prompted the development of aircraft and space facilities—in spite of their attendant costs and more limited availability.

8.3. Aircraft

Aircraft-based experiments provide longer durations for experiments without the impact loads seen in drop towers. However, aircraft are normally not able to execute adequately precise free-fall trajectories, and buoyant motion and other accelerative disturbances are usually not entirely suppressed. Thus, while aircraft experimentation is a valuable source of low-gravity test time, the test results generally only indicate trends with reduced buoyancy, and are not representative of true μg behavior. With the exception of a multiuser chamber developed by the European Space Agency, aircraft-based combustion experiments have generally been performed in test apparatus tailored to single programs. See Lekan²¹⁶ for a description of aircraft μg facilities in the United States.

Perhaps because of the cost of operations and of designing man-rated experimental hardware for the aircraft, these facilities are just beginning to be exploited by combustion experimenters. While fundamental studies of all types may benefit from the extended test time available in aircraft, a major use involves the development of diagnostic techniques in order to extend initial work being done in the drop towers. The PIV method of quantifying two-dimensional flow fields can be enhanced by the introduction of intensified detector arrays for the low-g loads of aircraft, so that higher velocity flows can be observed. The intensified array cameras will also provide improved visualization of dim, near-limit flames, as well as potential for measurements of species concentrations when fitted with bandpass filters.

Another aspect of aircraft-based facilities that has been used to some extent by combustion workers is the ability to free-float experiment payloads within the cabin volume, and thereby reduce the ambient accelerations by at least an additional order of magnitude. Unfortunately, this advantage has been limited by problems of free-floating test containers striking objects during free flight due to the limited space available in the aircraft cabins. If this problem can

be resolved, the aircraft facilities could provide an effective test platform for fundamental combustion studies. For the present, however, these facilities are best suited for the development of experimental techniques to be used during space-based tests. In particular, access to these facilities is reasonably good with each flight providing up to 40 test periods of 15–20 s duration.

8.4. *Sounding Rockets*

Sounding rockets can provide 200–900 s test times at low gravity, ca. 10^{-4} g, which is sufficient to achieve steady-state conditions for most combustion systems.²¹⁷ Facilities of this type have been exploited by the European Space Agency and are just coming into use in the United States for combustion experiments, with two experiments involving flame spread across liquids and solids, respectively, currently under development at NASA Lewis. Available test volumes are in the range 0.5–0.8 m³ with payloads of 200–300 kg; however, generally only a portion of the space and weight is allocated to any one user. The available test period with sounding rockets is attractive but the experiment must withstand significant g forces (10–40 g) during takeoff and landing which places limitations on available instrumentation. There is also concern that acceleration levels will be impulsively raised by the rate control systems of the spin-stabilized rockets. Thus, while development of sounding rockets is to be encouraged as a possibly less expensive and more accessible facility than spacecraft, the value of this approach for μ g combustion experiments has not been established at this time.

8.5. *Spacecraft*

Many combustion processes demonstrate the most interesting effects when the propagation velocities are at their lowest. This often occurs near the limit of flammability in terms of either fuel–air or oxygen–inert gas ratio. While ground-based experimenters have been creative in identifying systems that propagate quickly enough to observe low-gravity behavior in very short duration tests, most combustion processes will not be fully explored without the longer test times of low Earth orbit.

Condensed-phase fuels are limited during ground-based tests to very thin or shallow fuels, because conducted heat into the depth of the fuel retards the progress of the spreading flame. Consequently, experiments directed toward understanding the spreading of flames over thermally thick fuels, which represent nearly all engineering materials, cannot be attempted under any near-limit conditions in the ground-based facilities. While a series of engineering tests were conducted during the 1973–1974 Skylab era,²¹⁸ the first flight experiments to provide extended duration

flame spreading data compatible with modeling efforts have involved several solid-surface combustion experiments during recent Space Shuttle missions. The apparatus involves a quiescent chamber having a volume of 40 L. The test samples consist of thermally thin ashless filter paper and thermally thick polymethylmethacrylate (PMMA), ignited by a hot-wire filament coated with nitrocellulose, in selected gaseous environments. The facility is instrumented with two cameras to provide side and top views of the combustion process, thermocouples on and near the sample to measure solid and gas temperatures, and temperature and pressure sensors to monitor the chamber environment. The results are being used to evaluate theories emphasizing the mechanism of flame propagation.^{153,161–163}

A second series of experiments on the space shuttle were flown in connection with the United States Microgravity Laboratory (USML) in July 1992 (USML-1), using a glove box facility. This arrangement provides a working and storage volume of 25 L and allows considerable interaction between the experiment operator and the apparatus using two glove ports. Instrumentation consists of video and film visualization, and pressure, temperature and gas composition sensors. Three combustion experiments were flown, as follows: smoldering combustion of polyurethane, the ignition and combustion of electrical wire insulation, and the combustion of candle flames. All three experiments were operated successfully but results at this stage are preliminary and further experience with the μ g environment will be required to fully exploit the glove box facility.

Other experiments are being developed for flight testing in the late 1990s, involving gaseous premixed and diffusion flames, and drop diffusion flames. Developing experiments of this type for the manned space environment, however, is a long, complex and costly process. Additionally, flight opportunities are limited so that each experiment involves substantial risk. Thus, much remains to be done to achieve a more or less conventional laboratory environment allowing full exploitation of the long duration μ g, and the experiment–operator interaction, of manned space flight. As a result, the potential of reduced-gravity environments to clarify fundamental concepts of combustion will remain incomplete for some time.

In summary, we feel that long-duration experiments at μ g offer great potential for developing a fundamental understanding of combustion phenomena, and the technology base needed for fire-safe spacecraft operations. To achieve this potential, however, efforts need to be made to increase access to space for combustion experiments, which could involve the use of sounding rockets, Space Shuttle facilities involving minimal crew interaction, and unmanned orbiting spacecraft. In this regard, recent interest in exploiting sounding rockets by various space agencies—worldwide—should provide an opportunity to assess the potential of this resource for

combustion experiments. The advantages of using such intermediate facilities with greater frequency would be to provide a means of developing combustion experiments for the unfamiliar μg environment, and thus reduce the risk and enhance the impact of the more complex and costly experiments using manned spacecraft.

9. CONCLUDING REMARKS

Historically, progress in combustion research can be divided into several periods. In the pre-1950s period, many of the basic combustion phenomena—deflagration and detonation waves, diffusion flames, flame quenching, flamefront instability, and chemical kinetic effects—were observed and attempts were made to describe them. The 1950s was a truly robust period when the theory of aerothermochemistry was formulated and systematic study of various combustion phenomena was undertaken by a dedicated group of researchers. Combustion became a separate scientific discipline during this period. The period from the early 1960s to the mid-1970s involved continued exploitation of this technology, with notable progress in environment and aerospace-related research. From the mid-1970s and throughout the 1980s, combustion research was unusually active. These intensified activities were motivated by concerns about energy efficiency and environmental quality, and facilitated by the appearance of the various 'tools' needed for rigorous study. These tools included activation-energy asymptotics for analytical studies, computational advances for numerical studies, and laser-based diagnostics for experimental studies.

While sophisticated tools have been developed for combustion research, truly significant progress has been hindered by the lack of 'clean' and well-defined combustion and flame phenomena, through which individual processes can be isolated and studied in depth. A major cause of difficulty has been buoyancy. It is therefore not unrealistic to anticipate that the current interest in μg combustion, if sustained, could usher in the fifth period of combustion research, during which many of the fundamental issues of combustion and fire safety are finally resolved in a rigorous manner. The extremely interesting μg results reported here provide a strong indication of the potential. In order to accomplish this goal, however, experimental investigations will have to be conducted nonintrusively, most likely via optical diagnostics. For certain experiments sufficient run time also is needed in order to achieve a steady state and to perform an adequate number of tests to achieve statistically significant results. These requirements suggest that many more combustion experiments will have to be conducted on board sounding rockets, the Space Shuttle and future space stations, in order to fully realize the potential benefits of μg combustion research.

Finally, while μg offers new opportunities for fundamental studies of combustion phenomena, there now is ample evidence that our current understanding of fire and explosion hazards at ng has questionable relevance at μg conditions. This is a serious matter due to the extraordinary value of spacecraft facilities and the high visibility of accidents in space. Thus, there is strong motivation for a comprehensive program of fire research for spacecraft environments, to avoid both fire-related accidents and overly stringent fire safety regulations that could excessively increase the cost of spacecraft operations and limit capabilities for the beneficial exploitation of space. Among the fire safety issues that need to be addressed, the possibility of providing fire-safe spacecraft environments deserves greater priority than in the past. In particular, there may be a potential to leave Earth-bound fire hazards behind in the controlled environment of spacecraft, which has substantial implications for manned space activities. The feasibility of this potential must be understood by wide-ranging interdisciplinary research involving the effects of various environments on flame properties as well as the health and performance of humans and other organisms in these environments.

Acknowledgements—It is a pleasure to acknowledge useful inputs from H. D. Ross and K. Sacksteder of the NASA Lewis Research Center, M. K. King of the NASA Headquarters, P. D. Ronney of the University of Southern California and M. Y. Bahadori of the SAI Corporation. The authors' research in areas discussed in this paper has been sponsored by the NASA Microgravity Science and Applications Division (through the Lewis Research Center), the National Science Foundation, the Department of Energy, the Army Research Office, the Air Force Office of Scientific Research, the Office of Naval Research and the National Institute of Standards and Technology. The U.S. Government is authorized to reproduce and distribute copies of this article for governmental purposes notwithstanding any copyright notation thereon.

REFERENCES

1. *Microgravity Combustion Science: A Program Overview*, NASA TM 101424 (1989).
2. *Microgravity Combustion Science: Progress, Plans and Opportunities*, NASA TM 105410 (1992).
3. LAW, C. K., *AIAA Paper No. 90-0120* (1990).
4. FAETH, G. M., *AIAA/IKI Microgravity Science Symposium, Moscow*, pp. 281–293, AIAA, Washington (1991).
5. SACKSTEDER, K., *23rd Symposium (International) on Combustion*, pp. 1589–1596, The Combustion Institute, Pittsburgh (1990).
6. POPE, S. B., *23rd Symposium (International) on Combustion*, pp. 591–612, The Combustion Institute, Pittsburgh (1990).
7. OSTRACH, S., *Adv. Heat Trans.* **8**, 161 (1972).
8. LAW, C. K., CHUNG, S. H. and SRINIVASAN, N., *Combust. Flame* **38**, 173 (1980).
9. KAWAKAMI, T., OKAJIMA, S. and INUMA, K., *23rd Symposium (International) on Combustion*, pp. 1663–1667, The Combustion Institute, Pittsburgh (1990).
10. WU, M.-S., KWON, S., DRISCOLL, J. F. and FAETH, G. M., *Combust. Sci. Technol.* **73**, 327 (1990).

11. WU, M.-S., KWON, S., DRISCOLL, J. F. and FAETH, G. M., *Combust. Sci. Technol.* **78**, 69 (1991).
12. KWON, S., WU, M.-S., DRISCOLL, J. F. and FAETH, G. M., *Combust. Flame* **88**, 221 (1992).
13. BRAY, K. N. C., *Turbulent Reacting Flows*, P. A. Libby and F. A. Williams (Eds), pp. 115–183, Springer-Verlag, Berlin (1980).
14. PETERS, N., *21st Symposium (International) on Combustion*, pp. 1231–1250, The Combustion Institute, Pittsburgh (1986).
15. LAW, C. K., *22nd Symposium (International) on Combustion*, pp. 1381–1402, The Combustion Institute, Pittsburgh (1988).
16. ABRAHAM, J., WILLIAMS, F. A. and BRACCO, F. V., *SAE Paper No. 850345* (1985).
17. KARLOVITZ, B., DENNISTON, D. K., KNAPPCHAFER, D. H. and WELLS, F. E., *4th Symposium (International) on Combustion*, pp. 613–620, The Combustion Institute, Pittsburgh (1952).
18. CLAVIN, P., *Prog. Energy Combust. Sci.* **11**, 1 (1985).
19. TSENG, L.-K., and FAETH, G. M., *Combust. Flame*, submitted.
20. SIVASHINSKY, G. I., *Acta Astronautica* **4**, 1177 (1977).
21. MATALON, M. and MATKOWSKY, B. J., *J. Fluid Mech.* **124**, 239 (1982).
22. PELCE, P. and CLAVIN, P., *J. Fluid Mech.* **124**, 219 (1982).
23. CHUNG, S. H. and LAW, C. K., *Combust. Flame* **75**, 309 (1989).
24. WILLIAMS, F. A., *Combustion Theory*, 2nd edn, p. 135, Benjamin/Cummings, Menlo Park, CA (1985).
25. KWON, S., TSENG, L.-K. and FAETH, G. M., *Combust. Flame* **90**, 230 (1992).
26. TSENG, L.-K., ISHMAIL, M. and FAETH, G. M., *Combust. Flame* **95**, 410 (1993).
27. PALM-LEIS, A. and STREHLOW, R. A., *Combust. Flame* **13**, 111 (1969).
28. MARKSTEIN, G. H., *Non-Steady Flame Propagation*, p. 22, MacMillan, New York (1964).
29. FRISTROM, R. M., *Phys. Fluids* **8**, 273 (1965).
30. DESHAIES, B. and CAMBRAY, P., *Combust. Flame* **82**, 361 (1990).
31. TAYLOR, S. C., Ph.D. Thesis, University of Leeds, 1991; see also, Dowdy, D. R., Smith, D. B., Taylor, S. C. and Williams, A., *23rd Symposium (International) on Combustion*, pp. 325–332, The Combustion Institute, Pittsburgh (1990).
32. WU, C. K. and LAW, C. K., *20th Symposium (International) on Combustion*, pp. 1941–1949, The Combustion Institute, Pittsburgh (1984).
33. LAW, C. K., ZHU, D. L. and YU, G., *21st Symposium (International) on Combustion*, pp. 1419–1421, The Combustion Institute, Pittsburgh (1986).
34. ZHU, D. L., EGOLFOPOULOS, F. N. and LAW, C. K., *22nd Symposium (International) on Combustion*, pp. 1539–1545, The Combustion Institute, Pittsburgh (1988).
35. LAW, C. K., *Reduced Kinetic Mechanisms for Application in Combustion*, B. Rogg and N. Peters (Eds), pp. 15–26, Springer-Verlag, Berlin (1993).
36. YAMAOKA, I. and TSUJI, H., *20th Symposium (International) on Combustion*, pp. 1883–1892, The Combustion Institute, Pittsburgh (1984).
37. ANDREWS, G. E., and BRADLEY, D. *Combust. Flame* **20**, 77 (1973).
38. SEARBY, G. and QUINARD, J., *Combust. Flame* **82**, 298 (1990).
39. WARNATZ, J., *24th Symposium (International) on Combustion*, pp. 553–579, The Combustion Institute, Pittsburgh (1992).
40. ENG, J. A., ZHU, D. L. and LAW, C. K., *2nd Microgravity Combustion Workshop*, NASA CP 10113, pp. 177–182 (1993); see also, Eng, J. A., Law, C. K. and Zhu, D. L., *AIAA Paper No. 94-0571* (1994).
41. LEWIS, B. and VON ELBE, G., *Combustion, Flames and Explosions of Gases*, p. 315, Academic Press, New York (1961).
42. STREHLOW, R. A., *Combustion Fundamentals*, 2nd edn, p. 380, McGraw-Hill, New York (1984).
43. SIVASHINSKY, G. I., *Ann. Rev. Fluid Mech.* **15**, 179 (1983).
44. GROFF, E. G., *Combust. Flame* **48**, 51 (1982).
45. ZEL'DOVICH, Y. B., *The Theory of Combustion and Detonation*, Academy of Sciences, Moscow (1944).
46. BARENBLATT, G. I., ZEL'DOVICH, Y. B. and ISTRATOV, A. G., *Zh. Prikl. Mekh. Tekh. Fiz.* **4**, 21 (1962).
47. SIVASHINSKY, G. I., *Combust. Sci. Technol.* **15**, 137 (1977).
48. MANTON, J., VON ELBE, G. and LEWIS, B., *J. Chem. Phys.* **20**, 153 (1952).
49. BREGON, B., GORDON, A. S. and WILLIAMS, F. A., *Combust. Flame* **33**, 33 (1978).
50. RAKIB, Z. and SIVASHINSKY, G. I., *Combust. Sci. Technol.* **54**, 69 (1987).
51. BÖHM, G. and CLUSIUS, K., *Z. Naturforsch.* **A3**, 386 (1948).
52. KAILASANATH, K., PATNAIK, G. and ORAN, E. S., Paper No. IAF-88-354, *39th Congr. of Int. Astro. Fed.*, Bangalore (1988).
53. PATNAIK, G., KAILASANATH, K., LASKY, K. J. and ORAN, E. S., *22nd Symposium (International) on Combustion*, pp. 1517–1526, The Combustion Institute, Pittsburgh (1988).
54. PATNAIK, G. and KAILASANATH, K., *23rd Symposium (International) on Combustion*, pp. 1641–1647, The Combustion Institute, Pittsburgh (1990).
55. PATNAIK, G. and KAILASANATH, K., *AIAA Paper No. 91-0784* (1991).
56. PATNAIK, G. and KAILASANATH, K., *24th Symposium (International) on Combustion*, pp. 189–195, The Combustion Institute, Pittsburgh (1992).
57. KAILASANATH, K., PATNAIK, G. and ORAN, E. S., NASA Contractors' Report 191051 (1993).
58. KAILASANATH, K., GANGULY, K. and PATNAIK, G., *Prog. Aero. Astro.* **151**, 247 (1993).
59. STREHLOW, R. A., NOE, K. A. and WHERLEY, B. L., *21st Symposium (International) on Combustion*, pp. 1899–1908, The Combustion Institute, Pittsburgh (1986).
60. KEE, R. J., MILLER, J. A., EVANS, G. H. and DIXON-LEWIS, G., *22nd Symposium (International) on Combustion*, pp. 1479–1494, The Combustion Institute, Pittsburgh (1988).
61. DUNSKY, C. and FERNANDEZ-PELLO, A., *23rd Symposium (International) on Combustion*, pp. 1657–1662, The Combustion Institute, Pittsburgh (1990).
62. DUNSKY, C. M., *24th Symposium (International) on Combustion*, pp. 177–187, The Combustion Institute, Pittsburgh (1992).
63. DUROX, D., *24th Symposium (International) on Combustion*, pp. 197–204, The Combustion Institute, Pittsburgh (1992).
64. RONNEY, P. D. and WACHMAN, H. Y., *Combust. Flame* **62**, 107 (1985).
65. RONNEY, P. D., *Combust. Flame* **62**, 121 (1985).
66. RONNEY, P. D., *22nd Symposium (International) on Combustion*, pp. 1615–1623, The Combustion Institute, Pittsburgh (1988).
67. RONNEY, P. D., *Combust. Sci. Technol.* **59**, 123 (1988).
68. ABBUD-MADRID, A. and RONNEY, P. D., *23rd Symposium (International) on Combustion*, pp. 423–431, The Combustion Institute, Pittsburgh (1990).
69. WHALING, K. N., ABBUD-MADRID, A. and RONNEY, P. D., Fall Technical Meeting, Western States Section, The Combustion Institute, Pittsburgh (1990).

70. RONNEY, P. D., *Combust. Flame* **82**, 1 (1990).
71. RONNEY, P. D. and SIVASHINSKY, G. I., *SIAM J. Appl. Math.* **49**, 1029 (1989).
72. FARMER, J. R. and RONNEY, P. D., *Combust. Sci. Technol.* **73**, 555 (1990).
73. LAW, C. K. and EGOLFOPOULOS, F. N., *23rd Symposium (International) on Combustion*, pp. 413–421, The Combustion Institute, Pittsburgh (1990).
74. LAW, C. K. and EGOLFOPOULOS, F. N., *24th Symposium (International) on Combustion*, pp. 137–144, The Combustion Institute, Pittsburgh (1992).
75. BUCKMASTER, J. and WEERATUNGA, S., *Combust. Sci. Tech.* **35**, 287 (1984).
76. BUCKMASTER, J. D., JOULIN, G. and RONNEY, P. D., *Combust. Flame* **79**, 381 (1990); *Ibid.* **84**, 411 (1991).
77. BUCKMASTER, J., GESSMAN, R. and RONNEY, P., *24th Symposium (International) on Combustion*, pp. 53–59, The Combustion Institute, Pittsburgh (1992).
78. BUCKMASTER, J., RONNEY, P. D. and SMOOKE, M., *AIAA Paper No. 93-0712* (1993).
79. BUCKMASTER, J., SMOOKE, M. and GIOVANGIGLI, V., *Combust. Flame* **94**, 113 (1993).
80. HAWTHORNE, W. R., WEDDELL, D. S. and HOTTEL, H. C., *3rd Symposium on Combustion, Flame and Explosion Phenomena*, pp. 266–288, The Combustion Institute, Pittsburgh (1949).
81. BURKE, S. P. and SCHUMANN, T. E. W., *Ind. Eng. Chem.* **20**, 998 (1928).
82. BILGER, R. W., *Prog. Energy Combust. Sci.* **1**, 87 (1976).
83. BILGER, R. W., *Combust. Flame* **30**, 277 (1977).
84. GORE, J. P. and FAETH, G. M., *21st Symposium (International) on Combustion*, pp. 1521–1531, The Combustion Institute, Pittsburgh (1986).
85. GORE, J. P. and FAETH, G. M., *J. Heat Trans.* **110**, 173 (1988).
86. SIVATHANU, Y. R. and FAETH, G. M., *Combust. Flame* **82**, 150 (1990).
87. FAETH, G. M., GORE, J. P., CHUECH, S. G. and JENG, S.-M., *Ann. Rev. Num. Fluid Mech. Heat Trans.* **2**, 1 (1989).
88. LAW, C. K. and CHUNG, S. H., *Combust. Sci. Technol.* **29**, 129 (1982).
89. CHUNG, S. H. and LAW, C. K., *Combust. Flame* **52**, 59 (1983).
90. FAETH, G. M. and SAMUELSON, G. S., *Prog. Energy Combust. Sci.* **12**, 305 (1986).
91. BILGER, R. W., *22nd Symposium (International) on Combustion*, pp. 475–488, The Combustion Institute, Pittsburgh (1988).
92. SPALDING, D. B., *Combustion and Mass Transfer*, Chapter 10, Pergamon Press, New York (1979).
93. COCHRAN, T. H. and MASICA, W. J., *NASA TN D-5872* (1970).
94. COCHRAN, T. H. and MASICA, W. J., *13th Symposium (International) on Combustion*, pp. 821–829, The Combustion Institute, Pittsburgh (1970).
95. EDELMAN, R. B., FORTUNE, O. F., WEILERSTEIN, G., COCHRAN, T. H. and HAGGARD, J. B., Jr, *14th Symposium (International) on Combustion*, pp. 399–412, The Combustion Institute, Pittsburgh (1972).
96. HAGGARD, J. B., Jr and COCHRAN, T. H., *Combust. Sci. Technol.* **5**, 291 (1972).
97. HAGGARD, J. B., Jr, *NASA TN D-6523* (1972).
98. HAGGARD, J. B., Jr and COCHRAN, T. H., *NASA TN D-7165* (1973).
99. HAGGARD, J. B., Jr, *NASA TP1841* (1981).
100. KLAJN, M. and OPPENHEIM, A. K., *19th Symposium (International) on Combustion*, pp. 223–235, The Combustion Institute, Pittsburgh (1982).
101. EDELMAN, R. B. and BAHADORI, M. Y., *Acta Astronautica* **13**, 681 (1986).
102. EDELMAN, R. B., BAHADORI, M. Y., OLSON, S. L. and STOCKER, D. P., *AIAA Paper No. 88-0645* (1988).
103. BAHADORI, M. Y., STOCKER, D. P. and EDELMAN, R. B., *AIAA Paper No. 90-0651* (1990).
104. BAHADORI, M. Y., EDELMAN, R. B., STOCKER, D. P. and OLSON, S. L., *AIAA J.* **28**, 236 (1990).
105. BAHADORI, M. Y., EDELMAN, R. B., SOTOS, R. G. and STOCKER, D. P., *AIAA Paper No. 91-0719* (1991).
106. BAHADORI, M. Y., EDELMAN, R. B., STOCKER, D. P., SOTOS, R. G. and VAUGHAN, D. F., *AIAA Paper No. 92-0243* (1992).
107. BAHADORI, M. Y., STOCKER, D. P., VAUGHAN, D. F. and ZHOU, L., *2nd International Microgravity Combustion Workshop*, pp. 91–105, NASA CP 10113 (1993).
108. MORTAZAVI, S., SUNDERLAND, P. B., JURNG, J., KÖYLÜ, Ü. Ö. and FAETH, G. M., *AIAA Paper No. 93-0708* (1993).
109. KÖYLÜ, Ü. Ö., SUNDERLAND, P. B., MORTAZAVI, S. and FAETH, G. M., *AIAA Paper No. 94-0428* (1994).
110. ELLZEY, J. L. and ORAN, E. S., *23rd Symposium (International) on Combustion*, pp. 1635–1640, The Combustion Institute, Pittsburgh (1990).
111. BAHADORI, M. Y., VAUGHAN, D. F., STOCKER, D. P., WEILAND, K. J. and EDELMAN, R. B., *Spring Technical Meeting*, pp. 75–79, Central States Section, The Combustion Institute, Pittsburgh (1992).
112. VISKANTA, R. and MENGÜC, M. P., *Prog. Energy Combust. Sci.* **13**, 97 (1987).
113. TIEN, C. L. and LEE, S. C., *Prog. Energy Combust. Sci.* **8**, 41 (1982).
114. KÖYLÜ, Ü. Ö. and FAETH, G. M., *Combust. Flame* **87**, 61 (1991).
115. KÖYLÜ, Ü. Ö., SIVATHANU, Y. R. and FAETH, G. M., *3rd International Symposium on Fire Safety Science*, pp. 625–634, Elsevier, London (1991).
116. FRIEDMAN, R., *1st International Symposium on Fire Safety Science*, pp. 349–359, Hemisphere, Washington (1986).
117. GLASSMAN, I., *22nd Symposium (International) on Combustion*, pp. 295–311, The Combustion Institute, Pittsburgh (1988).
118. HAYNES, B. S. and WAGNER, H. G., *Combust. Sci. Technol.* **7**, 229 (1981).
119. SIVATHANU, Y. R. and FAETH, G. M., *Combust. Flame* **81**, 133 (1990).
120. SCHUG, K. P., MANHEIMER-TIMNAT, Y., YACCARINO, P. and GLASSMAN, I., *Combust. Sci. Technol.* **22**, 235 (1980).
121. SUNDERLAND, P. B., MORTAZAVI, S., FAETH, G. M. and URBAN, D. L., *Combust. Flame* **96**, 97 (1994).
122. FLOWER, W. L. and BOWMAN, C. T., *20th Symposium (International) on Combustion*, pp. 1035–1044, The Combustion Institute, Pittsburgh (1984).
123. FLOWER, W. L. and BOWMAN, C. T., *21st Symposium (International) on Combustion*, pp. 1115–1124, The Combustion Institute, Pittsburgh (1986).
124. LEE, J. H. S., PERALDI, O. and KNYSTAUTAS, R., *2nd International Microgravity Combustion Workshop*, pp. 189–195, NASA CP 10113 (1993).
125. ROSS, H. D., FACCA, L. T., BERLAD, A. and TANGIRALA, V., *NASA TM 101371* (1989).
126. BERLAD, A., ROSS, H., FACCA, L. and TANGIRALA, V., *AIAA Paper 89-0500* (1989).
127. BERLAD, A. L., ROSS, H., FACCA, L. and TANGIRALA, V., *Combust. Flame* **82**, 448 (1990).
128. BERLAD, A. L., ROSS, H., FACCA, L. and TANGIRALA, V., *J. Prop. Power* **7**, 5 (1991).
129. BERLAD, A. L. and TANGIRALA, V., *Prog. Astron. Aero.* **132**, 59 (1991).
130. KUMAR, R. K., BOWLES, E. M. and MINTZ, K. J., *Combust. Flame* **89**, 320 (1992).
131. SESHADRI, K., BERLAD, A. L. and TANGIRALA, V., *Combust. Flame* **89**, 333 (1992).

132. RAYLEIGH, LORD, *The Theory of Sound*, Dover (1945).
133. HEGDE, U., ROSS, H. D. and FACCA, L. T., *Combust. Sci. Technol.* **94**, 279 (1993).
134. BUCKMASTER, J. and CLAVIN, P., *24th Symposium (International) on Combustion*, pp. 29–36, The Combustion Institute, Pittsburgh (1992).
135. DOSANJH, S., PETERSON, J., FERNANDEZ-PELLO, A. C. and PAGNI, P. J., *Acta Astronautica* **13**, 689 (1986).
136. DOSANJH, S. S., PAGNI, P. J. and FERNANDEZ-PELLO, A. C., *Combust. Flame* **68**, 131 (1987).
137. TORERO, J. L., FERNANDEZ-PELLO, A. C. and URBAN, D., *AIAA Paper No. 93-0829* (1993).
138. FERNANDEZ-PELLO, A. C. and PAGNI, P. J., *2nd International Microgravity Combustion Workshop*, pp. 251–256, NASA CP 10113 (1993).
139. HOWARD, J. B., *2nd International Microgravity Combustion Workshop*, pp. 73–79, NASA CP 10113 (1993).
140. MUNIR, Z. A. and ANSELM-TAMBURI, U., *Mater. Sci. Rep.* **3**, 277 (1989).
141. HOLT, J. B. and DUNMEAD, S. D., *Annu. Rev. Mater. Sci.* **21**, 305 (1991).
142. VARMA, A. and LEBRAT, J.-P., *Chem. Eng. Sci.* **47**, 2179 (1992).
143. GLASSMAN, I., BREZINSKY, K. and LAW, C. K., *Combustion Synthesis of Materials in Microgravity*, proposal to NASA (1991).
144. MAKINO, A. and LAW, C. K., *24th Symposium (International) on Combustion*, pp. 1883–1891, The Combustion Institute, Pittsburgh (1992).
145. MOORE, J. J., *AIAA Paper No. 93-0830* (1993).
146. FERNANDEZ-PELLO, A. C. and HIRANO, T., *Combust. Sci. Technol.* **32**, 1 (1983).
147. DI BLASI, C., CRESCITELLI, S. and RUSSO, G., *Combust. Flame* **72**, 205 (1988).
148. WEST, J., BHATTACHARJEE, S. and ALTENKIRCH, R. A., *Fire and Combustion Systems*, HTD-Vol. 199, A. M. Kanury and M. Q. Brewster (Eds), pp. 151–157, ASME, New York (1992).
149. FERKUL, P. V. and T'EN, J. S., *AIAA Paper No. 93-0827* (1993).
150. ZHOU, L. and FERNANDEZ-PELLO, A. C., *24th Symposium (International) on Combustion*, pp. 1721–1728, The Combustion Institute, Pittsburgh (1992).
151. OLSON, S. L., FERKUL, P. V. and T'EN, J. S., *22nd Symposium (International) on Combustion*, pp. 1213–1221, The Combustion Institute, Pittsburgh (1988).
152. OLSON, S. L., *Combust. Sci. Technol.* **76**, 233 (1991).
153. ALTENKIRCH, R. A., BHATTACHARJEE, S., OLSON, S. L. and WEST, J., *IKI/AIAA Microgravity Science Symposium, Moscow*, pp. 305–313, AIAA, Washington (1991).
154. BHATTACHARJEE, S. and ALTENKIRCH, R. A., *24th Symposium (International) on Combustion*, pp. 1669–1676, The Combustion Institute, Pittsburgh (1992).
155. BONNEAU, L., JOULAIN, P., MOST, J. M. and FERNANDEZ-PELLO, A. C., *AIAA Paper No. 93-0826* (1993).
156. T'EN, J. S., *Combust. Flame* **65**, 31 (1986).
157. FOUTCH, D. W. and T'EN, J. S., *AIAA J.* **25**, 972 (1987).
158. T'EN, J. S., *Combust. Flame* **80**, 355 (1990).
159. CHAO, B. H., LAW, C. K. and T'EN, J. S., *23rd Symposium (International) on Combustion*, pp. 9–18, The Combustion Institute, Pittsburgh (1990).
160. CHAO, B. H., LAW, C. K., *Combust. Flame* **92**, 1 (1993).
161. BHATTACHARJEE, S., ALTENKIRCH, R. A., SRIKANTARAH, N. and VEDHA-NAYAGAM, M., *Combust. Sci. Technol.* **69**, 1 (1990).
162. ALTENKIRCH, R. A. and BHATTACHARJEE, S., *AIAA Prog. Astro. Aero.* **130**, 723 (1990).
163. BHATTACHARJEE, S. and ALTENKIRCH, R. A., *23rd Symposium (International) on Combustion*, pp. 1627–1633, The Combustion Institute, Pittsburgh (1990).
164. SIRIGNANO, W. A., *Combust. Sci. Technol.* **6**, 95 (1972).
165. ROSS, H. D. and SOTOS, R. G., *23rd Symposium (International) on Combustion*, pp. 1649–1655, The Combustion Institute, Pittsburgh (1990).
166. MILLER, F. J. and ROSS, H., *24th Symposium (International) on Combustion*, pp. 1703–1711, The Combustion Institute, Pittsburgh (1992).
167. SCHILLER, D. N. and SIRIGNANO, W. A., *AIAA Paper No. 93-0825* (1993).
168. LAW, C. K., *Prog. Energy Combust. Sci.* **8**, 171, 1982.
169. KUMAGAI, S. and ISODA, H., *6th Symposium (International) on Combustion*, pp. 726–731, The Combustion Institute, Pittsburgh (1956).
170. KUMAGAI, S., SAKAI, T. and OKAJIMA, S., *13th Symposium (International) on Combustion*, pp. 779–785, The Combustion Institute, Pittsburgh (1970).
171. CHAUVEAU, C. and GÖKALP, I., *7th European Symposium on Materials and Fluid Sciences in Microgravity* (1989).
172. HARA, H. and KUMAGAI, S., *23rd Symposium (International) on Combustion*, pp. 1605–1610, The Combustion Institute, Pittsburgh (1990).
173. MATALON, M. and LAW, C. K., *Combust. Flame* **50**, 219 (1983); *Ibid*, **59**, 213 (1985).
174. LAZAR, R. S. and FAETH, G. M., *13th Symposium (International) on Combustion*, pp. 801–811, The Combustion Institute, Pittsburgh (1970).
175. FAETH, G. M., DOMINICIS, D. P., TULPINSKY, J. F. and OLSON, D. R., *12th Symposium (International) on Combustion*, pp. 9–18, The Combustion Institute, Pittsburgh (1968).
176. SPALDING, D. B., *ARS J.* **29**, 828 (1959).
177. ROSNER, D. E., *AIAA J.* **5**, 163 (1967).
178. SHUEN, J. S., YANG, V. and HSIAO, C. C., *Combust. Flame* **89**, 286 (1992).
179. FAETH, G. M., *Prog. Energy Combust. Sci.* **3**, 191 (1977).
180. SHAW, B. D., DRYER, F. L., WILLIAMS, F. A. and HAGGARD, J. B., *Acta Astronautica* **17**, 1195 (1988).
181. CHOI, M. Y., DRYER, F. L. and HAGGARD, J. B., Jr, *23rd Symposium (International) on Combustion*, pp. 1597–1604, The Combustion Institute, Pittsburgh (1990).
182. CHOI, M. Y., DRYER, F. L., GREEN, G. J. and SANGIOVANNI, J. J., *AIAA Paper No. 93-0823* (1993).
183. RANDOLPH, A. L. and LAW, C. K., *Combust. Flame* **64**, 267 (1986).
184. JACKSON, G. and AVEDISIAN, C. T., *AIAA Paper 93-0130* (1993).
185. WILLIAMS, F. A., *Prog. Aero. Astro.* **73**, Chap. 2 (1981).
186. CHUNG, S. H. and LAW, C. K., *Combust. Flame* **64**, 237 (1986).
187. CHO, S. Y., CHOI, M. Y. and DRYER, F. L., *23rd Symposium (International) on Combustion*, pp. 1611–1617, The Combustion Institute, Pittsburgh (1990).
188. CHO, S. Y., YETTER, R. A. and DRYER, F. L., *J. Comp. Phys.* **102**, 160 (1992).
189. YANG, J. C., JACKSON, G. S. and AVEDISIAN, C. T., *23rd Symposium (International) on Combustion*, pp. 1619–1625, The Combustion Institute, Pittsburgh (1990).
190. LEE, A. and LAW, C. K., *Combust. Sci. Technol.* **86**, 253 (1992).
191. CHOI, M. Y., CHO, S. Y., STEIN, Y. S. and DRYER, F. L., Fall Technical Meeting of the Eastern States Section, The Combustion Institute, Pittsburgh (1990).
192. FAETH, G. M. and OLSON, D. R., *SAE Trans.* 1793 (1968).
193. LAW, C. K., *AIChE J.* **24**, 626 (1978).
194. MAKINO, A. and LAW, C. K., *Combust. Flame* **73**, 331 (1988).

195. WANG, C. H., LIU, X. Q. and LAW, C. K., *Combust. Flame* **56**, 175 (1984).
196. LAW, C. K. and LAW, H. K., *Modern Developments in Energy, Combustion and Spectroscopy*, F. A. Williams, A. K. Oppenheim, D. B. Olfe and M. Lapp (Eds), Pergamon Press, New York, in press.
197. JACKSON, G. S. and AVEDISIAN, C. T., *2nd International Microgravity Combustion Workshop*, pp. 331-337, NASA CP 10113 (1993).
198. WANG, C. H. and LAW, C. K., *Combust. Flame* **59**, 53 (1985).
199. YANG, J. C., JACKSON, G. S. and AVEDISIAN, C. T., *23rd Symposium (International) on Combustion*, pp. 1619-1625, The Combustion Institute, Pittsburgh (1990).
200. CARLETON, F. B. and WEINBERG, F. J., *Nature* **330**, 635 (1987).
201. ROSS, H. D., SOTOS, R. G. and T'EN, J. S., *Combust. Sci. Technol.* **75**, 155 (1991).
202. STEINBERG, T. A., WILSON, D. B. and BENZ, F., *Combust. Flame* **88**, 309 (1992); *Ibid.* **91**, 200 (1992).
203. SPUNKLER, C. M., NASA TM 81728 (1981).
204. ROSS, H. D., *2nd Microgravity Combustion Workshop*, pp. 9-19, NASA CP 10113 (1993).
205. HUGGETT, C., *Aerosp. Med.* **40**, 1176 (1969).
206. HUGGETT, C., *Combust. Flame* **20**, 140 (1973).
207. FRIEDMAN, R. and SACKSTEDER, K. R., NASA TM 100944 (1988).
208. CARHART, H. W., *Spacecraft Fire Safety*, J. M. Margle (Ed.), pp. 51-57, NASA CP-2476 (1987).
209. MCALEVEY, R. F., III and MAGEE, R. S., *J. Spacecraft Rockets* **4**, 1390 (1967).
210. GANN, R. G., STONE, J. P., TATEM, P. L., WILLIAMS, F. W. and CARHART, H. W., *Combust. Sci. Technol.* **18**, 155 (1978).
211. HERRIGAN, D. J., *The Physiological Basis for Spacecraft Environmental Limits*, J. M. Waligora (Ed.), pp. 1-15, NASA RP-1045 (1979).
212. KNIGHT, D. R., *Spacecraft Fire Safety*, J. M. Margle (Ed.), pp. 59-64, NASA CP-2476 (1987).
213. WILLENBERG, H. J. *AIAA/IKI Microgravity Science Symposium, Moscow*, pp. 317-323, AIAA, Washington (1991).
214. YOUNGBLOOD, W. W. and VEDHA-NAYAGAM, M., NASA CR 185147 (1989).
215. DE RIS, J., *Spacecraft Fire Safety*, J. M. Margle (Ed.), pp. 43-49, NASA CP-2476 (1987).
216. LEKAN, J., NASA TM 101397 (1989); also *AIAA Paper No. 89-0236* (1989).
217. WESSLING, F. C. and MAYBEE, G. W., *AIAA/IKI Microgravity Science Symposium, Moscow*, pp. 356-366, AIAA, Washington (1991).
218. KIMZEY, J. H., *Proceedings of the Third Space Processing Symposium on Skylab Results*, Vol. 1, pp. 115-130, NASA TM X-70252 (1974).

SHORT COMMUNICATION

Extinction of Premixed Methane-Air Flames With Reduced Reaction Mechanism

J. K. BECHTOLD and C. K. LAW *Department of Mechanical and Aerospace
Engineering, Princeton University, Princeton, New Jersey 08544*

(Received December 15, 1993; in final form May 2, 1994)

ABSTRACT—We examine the extinction characteristics of premixed methane-air flames subject to volumetric heat loss with a reduced reaction mechanism. A combination of rate-ratio and large activation energy asymptotics is employed to resolve the multi-layered flame structure, and an expression for the non-adiabatic burning rate is obtained. Extinction turning points are reproduced and extinction conditions are found explicitly as a function of the various kinetic parameters. In particular the flame is found to extinguish when the burning rate (scaled by its adiabatic value) is reduced to a *nearly* constant value, roughly 0.6, over the entire range of system parameters, in agreement with previous asymptotic and numerical studies. The reduction in flame temperature at extinction is found to be less than that predicted by one-step models.

INTRODUCTION

In a recent study (Bechtold and Law, 1994a) we analyzed the basic structure of lean and stoichiometric premixed methane-air flames, and the dependence of the adiabatic burning rate on the dominant kinetic parameters was determined. The four-step C_1 -chain mechanism of Peters (1985) was adopted, and we employed a combination of rate-ratio and large activation energy asymptotics to resolve the flame structure. In particular, we retained the exponential temperature dependence of the reaction rates in the fuel consumption zone, where large activation energy asymptotics were employed. Thus our approach differs from that of several recent studies (cf. Peters and Williams, 1987; Seshadri and Peters, 1990) that have used rate-ratio techniques in all the layers within the flame structure. Results obtained from the two approaches were found to compare favorably, and we commented that one advantage of our approach is that, by retaining the exponential temperature dependence of the reaction rate terms, we are able to systematically study the response of premixed methane-air flames to small perturbations. The purpose of this note is to illustrate that point by examining the response of these flames to small volumetric heat loss.

A theory of flame extinction due to volumetric heat loss was first given by Spalding (1957) for a simple one-step kinetic scheme. Subsequent analyses using large activation energy asymptotics and a single overall reaction (Buckmaster, 1976; Joulin and Clavin, 1979), as well as a two-step mechanism (Chao and Law, 1994) have shown that, at the extinction limit, the burning rate is reduced to $e^{-1/2} \sim .61$ of its adiabatic value. This result is insensitive to the nature of the heat-loss, and is in good agreement with both experimental measurements (cf. Ronney, 1988) and recent numerical studies of non-steady flames using a one-step overall reaction (Sibulkin and Frendi, 1990). Numerical

studies with detailed kinetic mechanisms (Lakshmisha *et al.*, 1991; Law and Egolfopoulos, 1992) also predict that the burning rate will be reduced to about 60% of its adiabatic value at extinction. These results suggest a general condition for extinction that is insensitive to both the nature of the heat-loss and the reaction mechanism.

Kennel *et al.* (1991) have studied the extinction characteristics of lean hydrocarbon-air flames using Peters' (1985) four-step mechanism, modified for propane. They employed rate-ratio asymptotic techniques, which assume all reaction rates to have a power law dependence on temperature and use the ratio of individual reaction rates as the small parameter to analyze the various layers within the flame structure. However, the power-law dependence does not exhibit the same sensitivity to temperature variations as the Arrhenius approximation. Consequently, the computed burning velocities do not vary significantly with temperature perturbations, and the introduction of small heat losses into the system does not dramatically alter the results. Therefore Kennel *et al.* (1991) first employed rate-ratio asymptotics to derive an effective overall activation energy by fitting calculated values of burning velocity to an Arrhenius form, and then carried out an analysis including heat loss for a one-step reaction with the effective overall activation energy.

In the present study we use the same multi-step mechanism, but we employ activation energy asymptotics in the fuel consumption zone. We demonstrate how extinction of lean and stoichiometric premixed methane-air flames can be described in a systematic way using complex chemistry without having to calculate an effective overall activation energy for a subsequent one-step analysis. Thus extinction conditions are found explicitly in terms of the various physical parameters.

SUMMARY OF ASYMPTOTIC ANALYSIS

The multi-layered flame structure considered in the present study is identical to that analyzed by Peters and Williams (1987) and Bechtold and Law (1994a,b), which should be consulted for notation and details. The four-step mechanism derived by Peters (1985) is used, although it is further reduced to two steps by assuming steady state for the H -radical and partial equilibrium of the water-gas shift. Here we will only summarize the results of the analysis.

A formula for the burning rate is obtained as an eigenvalue of the structure problem in the fuel consumption zone. This zone has width $\delta \ll 1$ which is inversely proportional to the relevant activation energy. In terms of an appropriately stretched inner coordinate $\xi = \delta x$ the equation for the local fuel concentration y can be written as

$$\frac{d^2 y}{d\xi^2} = De^\theta y \sqrt{1-y} \exp[-\gamma(y+m\xi)], \quad (1)$$

$$y = -\xi \text{ at } \xi = -1, \text{ and } y = 0 \text{ as } \xi \rightarrow \infty. \quad (2)$$

The term $\sqrt{1-y}$ represents the local concentration of H , and a separate analysis is needed near the upstream location $\xi = -1$ to demonstrate the smooth transition to zero of H -atom concentration. That analysis also provides the upstream boundary condition (2). The parameter m can be fixed at $m = -2$ for methane-air flames (Peters

and Williams, 1987), and the two remaining parameters are lengthy expressions that depend on all the physico-chemical parameters in the problem. Finally, we note that the exponential terms in Equation (1) represent the temperature perturbation, where in particular θ is the reduction in flame temperature. These terms would be absent if rate-ratio asymptotics were employed in this region, and by setting γ and θ to zero we recover the structure problem first derived by Peters and Williams (1987).

In general the above system must be solved numerically, and the adiabatic burning rate $(\rho v)_{ad}$, corresponding to $\theta = 0$, was discussed in our previous paper (Bechtold and Law, 1994a). In the present paper our interest is in a qualitative description of the burning rate of nonadiabatic methane-air flames. Thus we avoid writing out the explicit expressions for De^θ and γ , and we simply note their dependence on ρv as

$$De^\theta \propto (\rho v)^{-4/5} \exp[-L/(\rho v)^2], \gamma \propto 1, \text{ lean mixture,} \quad (3)$$

$$De^\theta \propto \exp[-L/(\rho v)^2], \gamma \propto \sqrt{\rho v}, \text{ stoichiometric mixture.} \quad (4)$$

The different dependence on ρv in the two cases is due to the fact that these parameters depend on the local concentrations of various species, which in turn are proportional to the width of the oxidation layer. This layer was found to vary differently with the burning rate depending on whether the mixture is lean or stoichiometric.

RESULTS

For the lean mixture γ is seen to be independent of ρv . Therefore integration of the above system determines D , and hence the burning rate, explicitly as a function of γ , i.e. $De^\theta = \Gamma(\gamma)$. An expression for the nonadiabatic burning rate in terms of its adiabatic value follows immediately. In particular, we define the scaled burning rate $f = \rho v/(\rho v)_{ad}$ and the new heat loss parameter $\mathcal{H} = -f^2 \theta$ such that \mathcal{H} is independent of ρv . Thus our equation for f becomes

$$f^{4/5} \exp(\mathcal{H}/f^2) = 1. \quad (5)$$

This equation possesses the characteristic dual response curve for burning rate as a function of heat loss, as shown in Figure 1. The turning point, which we regard as the extinction point, occurs at $f_{ex} = e^{-1/2}$, $\mathcal{H}_{ex} = (2/5)e^{-1}$. This is precisely the value of f_{ex} obtained in previous studies using one-step kinetics (cf. Buckmaster, 1976; Joulin and Clavin, 1979). However, we note that the critical value of the heat-loss parameter is less than that predicted by the one-step model. At extinction, the drop in flame temperature due to heat loss is $\theta_{ex} = -2/5$, which in dimensional form can be expressed as

$$T_{ex} = T_b - \frac{2 R^0 T_b^2}{5 E}. \quad (6)$$

Thus, the multi-step scheme, which accounts for several reaction layers, predicts a reduction in flame temperature that is forty percent the value obtained in one-step models. This further emphasizes the importance of small, inherent heat losses on the extinction characteristics of real flames.

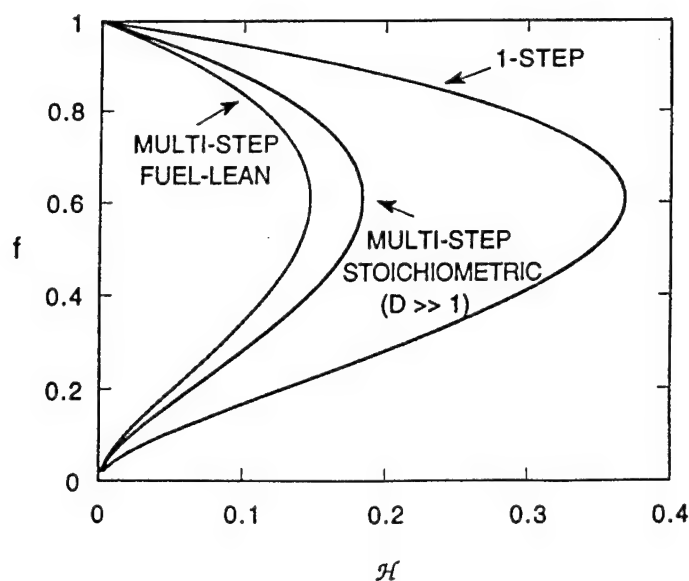


FIGURE 1 Plot of burning rate, scaled by the adiabatic value, as a function of heat loss, as found by both one-step and multi-step chemistry models.

For the stoichiometric mixture, γ is proportional to the square root of the burning rate, while the adiabatic Damköhler number D is independent of ρv . Response curves of γ as a function of D , determined by numerically solving Equations (1)–(2), are shown in Figure 2 for several values of the heat loss parameter $\mathcal{H} = -\gamma^4 \theta$.

By evaluating γ at the turning point, and comparing it to γ_{ad} for the corresponding value of D , we find that, at extinction, the burning rate is again reduced to approximate-

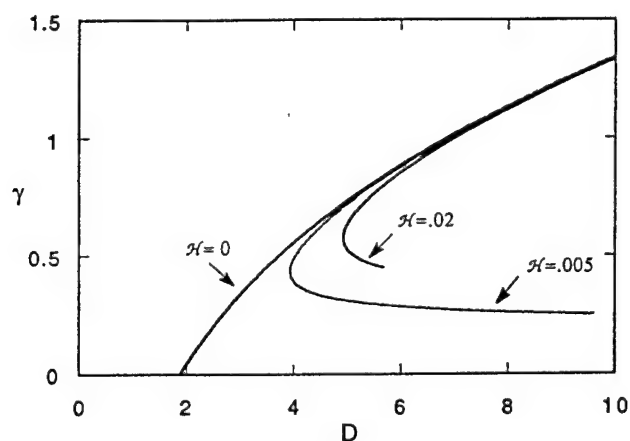


FIGURE 2 Plot of γ , representing the burning rate for stoichiometric flames, versus the parameter, D . Curves are found by solving equations (1) and (2) for three different values of the heat loss parameter, \mathcal{H} , with $m = -2$.

ly sixty percent of its adiabatic value over the entire range of D . For the stoichiometric mixture, it is not possible to write an explicit expression for the nonadiabatic burning rate in terms of the heat-loss parameter as was done for the fuel-lean case (i.e. Eq. (5)). However there are two limiting cases which yield such explicit relations. First, as demonstrated in Bechtold and Law (1994a), for $D \gg 1$ the structure becomes identical to the premixed burning regime of Liñán's (1974) diffusion flame analysis. Thus we can use his correlation function to obtain the relation

$$\gamma^2 \exp(\hat{\mathcal{H}}/\gamma^4) = 2D[0.6307m^2 - 1.344m + 1]^{-1} \quad (7)$$

In terms of the scaled burning rate f and the heat-loss parameter \mathcal{H} defined earlier this equation takes the form

$$f \exp(\mathcal{H}/f^2) = 1. \quad (8)$$

The turning point is now given by $f_{ex} = e^{-1/2}$, $\mathcal{H}_{ex} = e^{-1/2}$, and therefore the burning rate in this strongly-burning limit is again reduced to roughly .61 times its adiabatic value at extinction. Upon using the definition of \mathcal{H} we find T_{ex} in this limit to be

$$T_{ex} = T_b - \frac{R^0 T_b^2}{2E}, \quad (9)$$

and the reduction in flame temperature at extinction is one half that predicted by one-step models.

As D is decreased the flame slows down and the adiabatic burning rate reaches zero at $D = 15/8$. In this limit the structure reduces to that obtained using rate-ratio asymptotics (Peters and Williams, 1987), and the system (1)-(2) can be integrated to yield

$$D = \Gamma(\gamma_{ad}) \sim \frac{15}{8}(1 + \beta\gamma_{ad} + \dots), \quad (10)$$

where $\beta = (4/7 - m + 2mI)$ and $I = \int_0^1 \omega(1-\omega)\{1 + 2\omega + 3\omega^2 + \frac{3}{2}\omega^3\}^{1/2} d\omega = 0.291\dots$. The nonadiabatic burning rate, γ , in this limit is therefore obtained from

$$D \exp(\hat{\mathcal{H}}/\gamma^4) = \Gamma(\gamma). \quad (11)$$

The solution of this equation possesses a turning point at $\hat{\mathcal{H}}_{ex} = -\gamma^5 \Gamma'(\gamma)/4\Gamma(\gamma)$, and when this is inserted into Equation (11) we obtain the following equation for γ at extinction

$$D \exp(-\gamma_{ex} \Gamma'/4\Gamma) = \Gamma(\gamma_{ex}). \quad (12)$$

Upon expanding for small γ and making use of the relation (10) we obtain $\gamma_{ex} = \frac{4}{3}\gamma_{ad}$. Recalling that $\gamma \propto \sqrt{\rho v}$, this result implies that

$$(\rho v)_{ex} = \frac{16}{25}(\rho v)_{ad} = .64(\rho v)_{ad}. \quad (13)$$

This critical value, $f_{ex} = .64$, was also obtained for the lean hydrocarbon-air flames considered by Kennel *et al.* (1991) using a somewhat different approach. Thus, although f_{ex} is not constant over the entire range $15/8 < D < \infty$, it varies only slightly, i.e. $0.64 > f > 0.61$.

The value of the heat-loss needed to extinguish the flame, however, is found to decrease with D , eventually reaching zero as $D \rightarrow 15/8$. In particular, we find that $\mathcal{H}_{ex} \sim \sqrt{(\rho v)_{ad}}$ as this lower limit of D is approached. This is in contrast to the fuel-lean case, as well as previous asymptotic studies by Joulin and Clavin (1979) and Chao and Law (1994), that predict a constant value of the extinction heat-loss parameter over the entire range of system parameters. In this limit, the flame temperature at extinction is found to be

$$T_{ex} = T_b - \frac{R^0 T_b^2 \beta \gamma_{ad}}{E} \quad (14)$$

and therefore only a slight reduction in temperature is needed to cause extinction.

FURTHER DISCUSSIONS

We conclude our discussion of nonadiabatic methane-air flames by noting that phenomenological descriptions of extinction due to heat loss based on one-step chemistry, such as that given by Williams (1985), can be extended to include a broader class of multi-step kinetic mechanisms.

When a multi-step kinetic scheme is employed, the reaction zone can consist of several regions of varying widths in which species are continually produced and consumed. If the dominant reaction rate $\Omega(T)$ in one of the regions is extremely sensitive to temperature variations, then the burning rate can be expressed in terms of Ω as

$$\rho v \sim \sqrt{\int_{T_u}^{T_f} \Omega(T) dT} \quad (15)$$

For the particular case of an Arrhenius temperature dependence, this integral can be evaluated for large activation energy to yield

$$\rho v \sim \sqrt{A} \exp(-E/2R^0 T_f). \quad (16)$$

When small heat losses are present in the system, an energy balance reveals that the flame temperature will be lowered by a correspondingly small amount that is proportional to the inverse of the square of the burning rate, i.e. $T_f \sim T_{ad} - \delta/(\rho v)^2$, where $\delta \ll 1$ is the magnitude of the heat loss. This expression for T_f is inserted into Equation (16) to determine the nonadiabatic burning rate. For one-step chemistry with a single deficient reactant, the pre-exponential term A in Equation (16) is independent of ρv . Thus, the scaled burning rate $f = \rho v/(\rho v)_{ad}$ can be introduced to yield the familiar result

$$f \exp(\mathcal{L}/f^2) = 1, \quad (17)$$

if we assume $\delta = O(1/E)$, where \mathcal{L} is an appropriately chosen heat-loss parameter. The solution of this equation is known to possess a turning point at $f_{ex} = e^{-1/2}$.

In asymptotic analyses such as ours that adopt more detailed chemistry models the pre-exponential term in W contains contributions from the local concentrations of additional species in the flow field. These species can be consumed and produced in broader reaction zones that surround the temperature-sensitive region. The asymptotic studies have demonstrated that, in some cases, the species concentrations are proportional to the width of these broader layers, which in turn vary with burning rate. Therefore the reaction rate term may have a dependence on ρv of the form $\Omega \propto (\rho v)^\alpha$. When this is accounted for, the equation for the nonadiabatic burning rate f is modified to read

$$f^{(2-\alpha)/2} \exp(\mathcal{L}/f^2) = 1. \quad (18)$$

For all $\alpha < 2$, this equation has a turning point at $f_{ex} = e^{-1/2}$, $\mathcal{L}_{ex} = e^{-1}(2-\alpha)/4$. Thus, only the value of the heat-loss coefficient at extinction is affected.

It follows from Equation (18) that, for the particular case $\alpha = 2$, no expression is found for the adiabatic burning rate, and the above analysis is not appropriate. It is interesting to note that this situation corresponds to the stoichiometric methane-air flame in the limit $D \rightarrow 15/8$. As this lower limit of D is approached, the temperature perturbation becomes vanishingly small, and the rate-ratio asymptotic flame structure is recovered. Nevertheless, we have shown that the value of the burning rate at extinction is still reduced to slightly more than sixty percent of its adiabatic value.

ACKNOWLEDGEMENTS

This work was supported by the National Science Foundation and the Air Force Office of Scientific Research, under the technical monitoring of Dr. M. J. Linevsky and Dr. J. M. Tishkoff, respectively.

REFERENCES

- Bechtold, J. K. and Law, C. K. (1994a). The Structure of Premixed Methane-Air Flames with Large Activation Energy. *Combust. Flame*, **97**, 317.
- Bechtold, J. K. and Law, C. K. (1994b). Extinction of Premixed Methane-Air Flames with Volumetric Heat Loss. *Princeton University MAE Report No. T1998*.
- Buckmaster, J. D. (1976). The Quenching of Deflagration Waves. *Combust. Flame*, **26**, 151.
- Chao, B. H. and Law, C. K. (1994). Laminar Flame Propagation with Volumetric Heat Loss and Chain Branching-Terminating Reactions. *Int. J. Heat Mass Transfer*, **37**, 673.
- Joulin, G. and Clavin, P. (1979). Linear Stability Analysis of Nonadiabatic Flames: a Thermal-Diffusional Model. *Combust. Flame*, **35**, 139.
- Kennel, C., Göttgens, J. and Peters, N. (1991). The Basic Structure of Lean Propane Flames, *Twenty-Third Symposium (Int.) on Combustion*, The Combustion Institute, Pittsburgh, pp. 479-85.
- Lakshmisha, K. N., Paul, P. J. and Mukunda, H. S. (1991). On the Flammability Limit and Heat Loss in Flames with Detailed Chemistry, *Twenty-Third Symposium (Int.) on Combustion*, The Combustion Institute, Pittsburgh, pp. 433-440.
- Law, C. K. and Egolfopoulos, F. N. (1992). A Unified Chain-Loss Theory of Fundamental Flammability Limits, *Twenty-Fourth Symposium (Int.) on Combustion*, The Combustion Institute, Pittsburgh, pp. 137-144.
- Liñán, A. (1974). The Asymptotic Structure of Counterflow Diffusion Flames for Large Activation Energies. *Acta Astronautica*, **1**, 1007.
- Peters, N. (1985). Numerical Simulation of Combustion Phenomena, edited in R. Glowinski *et al.* (Ed.), *Lecture Notes in Physics*, **241**, Springer-Verlag, pp. 90-109.

- Peters, N. and Williams, F. A. (1987). The Asymptotic Structure of Stoichiometric Methane-Air Flames. *Combust. Flame*, **68**, 185.
- Ronney P. D. (1988). On the Mechanisms of Flame Propagation Limits and Extinguishment Processes at Microgravity, *Twenty-Second Symposium (Int.) on Combustion*, The Combustion Institute, Pittsburgh, pp. 1615-1623.
- Seshadri, K. and Peters, N. (1990). The Inner Structure of Methane-Air Flames. *Combust. Flame*, **81**, 96.
- Sibulkin, M. and Frendi, A. (1990). Prediction of Flammability Limit of an Unconfined Premixed Gas in the Absence of Gravity. *Combust. Flame*, **82**, 334.
- Spalding, D. B. (1957). A Theory of Inflammability Limits and Flame-Quenching. *Proc. Roy. Soc. London*, **A240**, 83.
- Williams, F. A. (1985). *Combustion Theory*, 2nd Ed., Addison-Wesley, Menlo Park CA.

EFFECTS OF PRESSURE AND DILUTION ON THE EXTINCTION OF COUNTERFLOW NONPREMIXED HYDROGEN-AIR FLAMES

P. PAPAS, I. GLASSMAN AND C. K. LAW

*Department of Mechanical and Aerospace Engineering
Princeton University Princeton, NJ 08544, USA*

To increase the understanding of the critical phenomena of ignition and extinction in high-speed propulsion devices, an experimental and computational study has been conducted on the strain-induced extinction of nonpremixed counterflow flames of diluted hydrogen against air. The study reports laser-Doppler velocimetry- (LDV-) determined local extinction strain rates of these flames, with various amounts of dilution and at pressures of 0.5 and 1.0 atm. The measured data compare well with results obtained from computational simulations with detailed chemistry and transport. Additional computational study on the effects of dilution and pressure shows that the extinction flame temperatures and strain rates exhibit a nonmonotonic variation with increasing pressure, which is characteristic of the explosion limits of homogeneous hydrogen-oxygen mixtures. This behavior is explained on the basis of the intrinsic chain branching-termination kinetics of hydrogen oxidation. The similarity in the dominant kinetic steps responsible for both the ignition/explosion and extinction phenomena is discussed.

Introduction

Recent interests in high-speed aeropropulsion have led to considerable research on hydrogen-oxygen chemistry [1,2] and its coupling to fluid flows. Because of the high-speed nature of the flow, the available residence time for mixing and chemical reactions is significantly reduced, leading to extreme difficulties in achieving ignition and preventing extinction. Consequently, studies of the ignition and extinction phenomena involving hydrogen-air mixtures are of importance to the development of high-speed aeropropulsion.

Recent studies on hydrogen-air flames have mostly been theoretical in nature, involving analyzing the structure and extinction limits of counterflow nonpremixed flames. Prominent contributions include Dixon-Lewis's detailed computational study [3] of the chemical flame structure and extinction limits in terms of the strain rate and oxygen concentrations, Gutheil and coworker's [4,5] computational and asymptotic analysis with emphasis on reduced mechanisms, and Darabiha and Candel's [6] computational study of the temperature effects on the ignition and extinction limits. Experimentally, Pellett and coworkers [7,8] have determined the extinction strain rates of diluted counterflow hydrogen-air flames based on global experimental parameters such as the average air velocity and nozzle diameter.

The present investigation aims to complement the previous studies along the following directions. Since the experimental extinction strain-rate data of Pellett et al. [7,8] are based on global parameters, there has

been substantial uncertainty in comparing these data with results from computational studies based on local strain rates. Thus, there exists the need for systematic, benchmark experimental data on hydrogen-air flame extinction based on local measurements. Consequently, local near-extinction strain rates of hydrogen-air counterflow flames with different extent of dilution, and at 0.5 and 1 atm pressure, were measured using laser-Doppler velocimetry (LDV). Comparisons of these experimental data with results obtained from computational simulations utilizing detailed transport and chemistry are then appropriate. When making these comparisons, the effect of the imposition of the various boundary conditions at the reactant stream exits in the computational simulation will be addressed.

Furthermore, computational simulations permit the study of the influence of pressure on the extinction states of hydrogen-oxygen flames. In this regard, it is to be noted that the combustion chamber within aeroengines will not only operate at different pressures, but the chamber pressure can also undergo strong fluctuations. Very few studies have been conducted on the effect of pressure on hydrogen-air flame extinction. Gutheil and Williams [5] have shown that the extinction strain rate increases with pressure. Because of the inherent chain nature of the hydrogen-oxygen chemistry, however, the explosion limits of homogeneous hydrogen-oxygen mixtures exhibit a nonmonotonic behavior with pressure, as shown in Fig. 4. Similar behavior has also been recently observed for the counterflow ignition of a hydrogen jet by a heated air jet [9]. Thus, it is of both

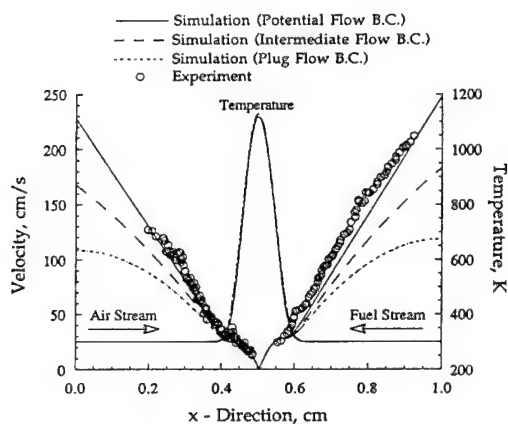


FIG. 1. Axial velocity profiles near extinction for a 16% hydrogen-84% nitrogen mixture impinging air at 1 atm. Reactant stream exit temperatures are 300 K.

fundamental and practical interest to explore if non-monotonic behavior also exists for flame extinction of counterflow nonpremixed flames.

In the next two sections, the experimental and computational aspects of this study are discussed and the results of the investigation reported.

Experimental

The extinction conditions of nonpremixed hydrogen-air flames were experimentally determined in a nozzle- or tube-generated counterflow configuration [10]. The fuel streams consisted of 12.0–18.5 mole% hydrogen in nitrogen, and the oxidizer stream was air. The diameters of the converging nozzles and tubes were between 0.7 and 1.4 cm. The separation distance between the nozzle and tube exits ranged from 0.6 to 1.4 cm. For each fuel concentration studied, extinction was accomplished by gradually increasing the velocities of both streams. Near-extinction strain rates were determined by LDV from axial velocity profiles obtained along the centerline [10]. In this study, the strain rate, K , is defined as half of the measured axial velocity gradient upstream of the flame on the air side. Typical experimental measurements of axial velocities near extinction are given in Fig. 1. The case depicted is for a 16% hydrogen mixture impinging onto an air stream at 1 atm pressure and room temperature, where x is the axial coordinate, with $x = 0$ and 1 cm corresponding to the air and fuel stream exits, respectively. The open circles in this figure represent the average of many dual-beam LDV measurements (usually 30) at a particular axial location along the stagnation streamline. The accuracy of these experimental measurements is within 10%, with measurements near the stagnation plane having a higher degree of inaccuracy. As will

be discussed, experimental axial velocity gradients were determined by fitting a straight line through the LDV measurements.

Computational

In order to allow for comparison with experimental data, numerical simulations were conducted using a quasi-one-dimensional opposed jet nonpremixed laminar flame model along the stagnation point streamline of the axis of symmetry. The corresponding steady-state mass, species, momentum, and energy conservation equations are solved using detailed chemical and transport schemes. The numerical code employed in this work is based on the scheme of Smooke and coworkers, which has been extensively described elsewhere [11]. Boundary conditions are applied at the air and fuel exits. The mass conservation that applies at the exit boundaries is $2b = -(du/dx)$, where u is the axial velocity. Conventionally, the outer frozen fields are assumed to be described by potential flow. The strain rate at the air boundary, $K = b_0$, then describes the constant axial velocity gradient. Another boundary condition commonly applied at the exit is that of plug flow ($b_0 = 0$), where the radial velocity is zero. Typically, the experimental boundary conditions of counterflow burners lie between those of potential and plug flows [10]. The boundary condition imposed in such cases would be determined from the experimental axial velocity gradient at the exits, namely, $b_0 = -(du/dx)_0/2$ and $b_1 = -(du/dx)_1/2$. The potential and plug flow boundary conditions may, therefore, be viewed as limiting cases [10]. The choice of boundary conditions for the outer frozen flow fields will be discussed further in the course of this paper.

The detailed hydrogen-oxygen reaction mechanism used consists of 19 elementary reaction steps with reverse reaction rates determined from equilibrium [1,2,12]. This mechanism is based on the scheme of Yetter et al. [1,2], which has been used to predict a variety of experimental data over a wide range of conditions [1,12]. The scheme utilized in this study, listed in Table 1, includes pressure falloff for the reaction $H_2O_2 + M = 2OH + M$. Thermochemical properties of the nine species in the reaction mechanism are also taken from Yetter et al. [2]. The molecular parameters used in the numerical simulation to evaluate the transport properties were obtained from the CHEMKIN data base [13].

Numerical simulations were conducted for various total pressures of a nitrogen-diluted hydrogen stream impinging an air stream. In particular, the steady-state condition just prior to extinction is examined. Figure 2 represents the variation of the maximum flame temperature of several mixtures as a function of the inverse of the air-side strain rate, $1/K$. The numerical simulation shown in Fig. 2 utilizes poten-

TABLE I
 Hydrogen/oxygen reaction mechanism^a

	A_f (mole cm ³ s)	n_f	E_{af} (KJ/mole)	T_{range} (K)	Reference
Chain reactions:					
1. $H + O_2 = OH + O$	1.92×10^{14}	0.00	68.78	962–2,577	15
2. $H_2 + O = OH + H$	5.08×10^{14}	2.67	26.33	297–2,495	16
3. $H_2 + OH = H + H_2O$	2.16×10^{18}	1.51	14.35	250–2,581	17
4. $OH + OH = H_2O + O$	1.23×10^{14}	2.62	–7.86	250–2,000	18
Formation and consumption of HO₂:					
5. $H + O_2 + M = HO_2 + M^c$	6.70×10^{19}	–1.42	0.00	200–2,000	19
6. $HO_2 + H = OH + OH$	1.69×10^{14}	0.00	3.66	298–773	20
7. $HO_2 + H = H_2 + O_2$	6.63×10^{13}	0.00	8.89	298–773	20
8. $HO_2 + OH = H_2O + O_2$	1.45×10^{16}	–1.00	0.00	298–1,400	20
9. $HO_2 + O = O_2 + OH$	1.81×10^{13}	0.00	–1.66	200–400	20
Recombination/dissociation reactions:					
10. $H_2 + M = H + H + M^c$	4.57×10^{19}	–1.40	436.80	600–2,000	20
11. $O + O + M = O_2 + M^c$	6.17×10^{15}	–0.50	0.00	2,000–10,000	20
12. $H + OH + M = H_2O + M^c$	2.25×10^{22}	–2.00	0.00	1,000–3,000	20
13. $O + H + M = OH + M^c$	4.72×10^{18}	–1.00	0.00		20
Formation and consumption of HOOH:					
14. $HO_2 + HO_2 = H_2O_2 + O_2$				300–1,100	21
	$k = 4.20 \times 10^{14} \exp(-50.13/RT) + 1.30E11 \exp(6.816/RT)$				
15. $H_2O_2 + M = OH + OH + M^{b,c}$					22
	$k_o = 1.20 \times 10^{17} \exp(-190.37/RT), F_c = 0.5$				
	$k_w = 2.95 \times 10^{14} \exp(-202.65/RT), N = 1.13$				
16. $H_2O_2 + OH = H_2O + HO_2$				250–1,250	23
	$k = 1.00 \times 10^{12} + 5.80 \times 10^{14} \exp(-39.99/RT)$				
17. $H_2O_2 + H = H_2 + HO_2$	4.82×10^{13}	0.00	33.25	283–800	20
18. $H_2O_2 + H = H_2O + OH$	1.00×10^{13}	0.00	15.02	283–800	24
19. $H_2O_2 + O = OH + HO_2$	9.55×10^{16}	2.00	16.61	250–800	20

^a $k = AT^n \exp(-E_a/RT)$; f denotes forward rate value.

^b $k_{15} = k_x \left[\frac{k_o/k_w}{1 + (k_o/k_w)} \right] F_c, X = \{1 + [\log(k_o/k_w)/N]^2\}^{-1}$.

^c $[M] = [N_2] + [O_2] + 2.5[H_2] + 12[H_2O] + [H] + [O] + [OH] + [H_2O_2] + [HO_2]$.

tial flow boundary conditions at the exits (300 K). It is seen that, for a given mixture and stream exit temperatures, as the inverse strain rate, $1/K$, is decreased, the maximum flame temperature decreases until the characteristic extinction turning point is exhibited. Extinction is then identified as the state of the turning point.

Results

Figure 3 shows the experimental and numerical density-weighted near-extinction strain rates, ρK_{ext} , plotted as a function of the fuel-stream hydrogen mole fraction, where ρ is the exit air density and K_{ext} is half the air axial velocity gradient. Overall, the ex-

perimentally and numerically determined near-extinction strain rates agree well. The only other experimental values for hydrogen-air extinction were reported by Pellett and coworkers [7,8]. Their value for the extinction strain rate of a 21% mole fraction hydrogen (79% nitrogen) mixture (622/s), shown in Fig. 3, was obtained with global experimental parameters, U_{air}/D_c , where U_{air} is the average air exit velocity and D_c represents an empirically selected weighting factor used to achieve agreement between opposed jet tubular and nozzle exit extinction results. Therefore, there is considerable uncertainty concerning this result.

Also shown in Fig. 3 are the computed results of Gutheil and Williams [5] and Darabiha and Candel [6], using the potential flow boundary condition. The

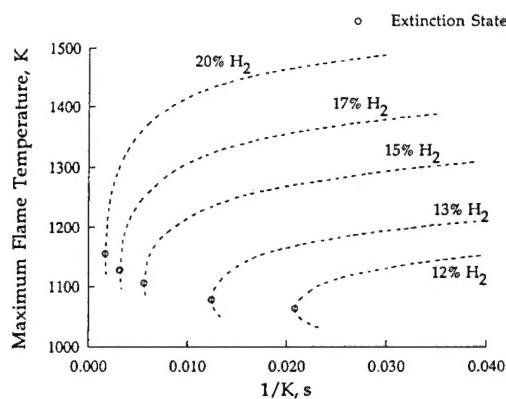


FIG. 2. Maximum flame temperature as a function of inverse strain rate for 20, 17, 15, 13, and 12% hydrogen-nitrogen mixtures impinging and an air stream at 1 atm. Reactant stream exit temperatures are 300 K.

extinction strain rates calculated by Gutheil and Williams [5] are in close agreement with the calculated results of this study. Darabiha and Candel's extinction strain rate for 16% mole fraction hydrogen in nitrogen, denoted by an arrow, is about 20% lower than the present value. Since the transport descriptions utilized by Darabiha and Candel and the current study are similar, the difference in the calculated extinction strain rates may be attributed to differences in the reaction schemes used.

At this point, it is necessary to examine the sensitivity of the results to the boundary conditions adopted. As stated previously, the experimental boundary conditions lie between those of potential

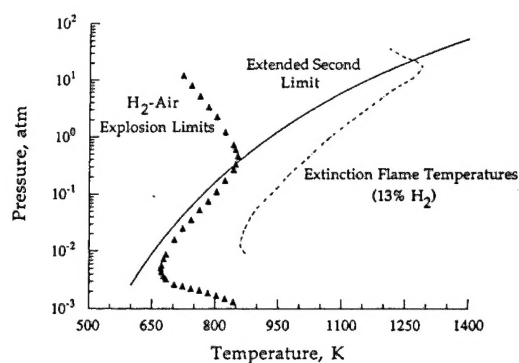


FIG. 4. Explosion limits for a stoichiometric hydrogen-air mixture, the "extended second limit," and extinction temperatures of a nonpremixed 13% hydrogen/87% nitrogen-air flame as a function of pressure.

and plug flows. Plotted in Fig. 1 are the near-extinction axial velocity profiles for a 16% hydrogen mixture issuing from a converging nozzle with diameter of 1.4 cm. The numerical simulations are the near-extinction axial velocity profiles using potential, plug, and an intermediate boundary condition. The intermediate boundary condition in Fig. 2 represents the extinction state for exit velocity values that are the average of the plug and potential flow cases. The strain rate defined from the experimental velocity profile over a region extending about 0.2 cm upstream from the reaction zone ($x \approx 0.4$ cm) is 200/s. Similarly, a strain rate of 245/s can be defined for the potential flow boundary condition. The extinction strain rates defined in the potential flow cases are

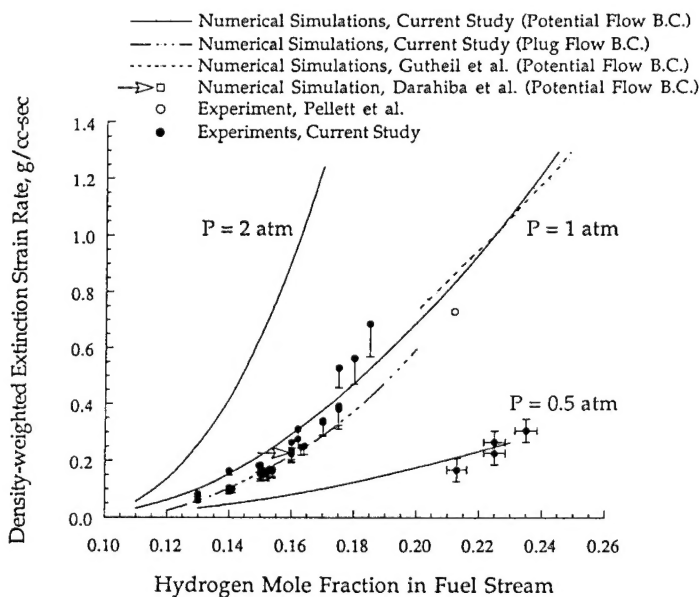


FIG. 3. Experimental and numerical values of air density-weighted extinction strain rates as a function of hydrogen mole fraction in nitrogen in the fuel stream. Reactant stream exit temperatures are 300 K. Unless specified otherwise, numerical simulations incorporate potential flow boundary conditions.

independent of the axial coordinate in the outer frozen flow field. Moreover, numerical simulations performed in this study have also demonstrated that the definition of the extinction strain rate for the potential flow is, to a great extent, also independent of the stagnation point location. For the plug and intermediate flows, however, the axial velocity gradient varies throughout the domain between the boundaries. Essentially, the strain rate defined for these cases just upstream of the reaction zone on the air side is represented by the maximum strain rate over the air-side axial domain. The maximum strain rate defined for the plug flow case of 185/s is 25% lower than the potential flow extinction strain rate. The flame temperature profiles near extinction of the numerical simulations for the case represented in Fig. 1, however, are largely independent of the type of boundary condition at the frozen reactant exits.

Maximum density-weighted extinction strain rates at 1 atm, obtained with plug flow boundary conditions, are plotted in Fig. 3 as a function of the hydrogen mole fraction in the fuel stream. The stagnation point locations for these plug flow cases are midway between the exits (0.5 cm). As expected, the experimental extinction strain-rate results of this study lie between those obtained with the potential and plug flow boundary conditions.

It is also important to realize that, similar to the computational results, the experimentally defined extinction strain rates also depend on several secondary factors and, therefore, are not unique. These factors include the axial velocity gradients at the nozzle exits, the location of the stagnation plane, and the region over which strain is defined. Consequently, the extinction strain rates reported here should be treated as approximate values. The experimental strain rates in Fig. 3 are based on local definitions obtained from least-squares fits of the experimental data immediately upstream of the reaction zone on the air side. These data have an uncertainty of 10% for the 12–16% hydrogen mixtures at 1 atm, 25% for the 17.0 and 18.5% mixtures at 1 atm, and 20% at 0.5 atm. Error bars for the 0.5-atm data also reflect uncertainties in the system pressure. In general, Fig. 3 shows that the experimentally determined strain rates for the dilute mixtures studied herein are characterized well by the numerically calculated values.

The effect of pressure on extinction of diluted hydrogen-air mixtures was also numerically investigated, using the potential flow boundary conditions for simplicity. Calculated maximum flame temperatures at extinction for a 13% hydrogen mixture are shown in Fig. 4 as a function of the system pressure. The temperature at both exits is 300 K. The relationship of extinction temperatures with pressure for this system exhibits the familiar nonmonotonic pressure dependency as observed for the homogeneous hydrogen-oxygen explosion limits. Specifically, the region to the right of this curve represents burning

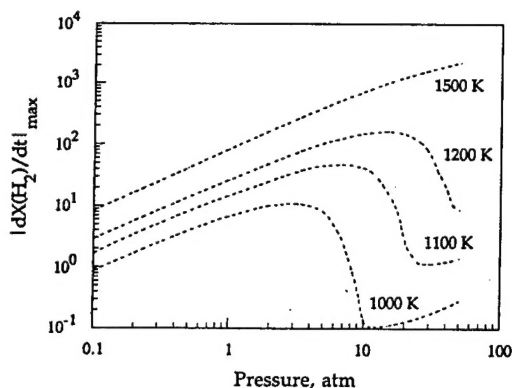


FIG. 5. Maximum reaction rate as a function of pressure for a 1% hydrogen, 0.5% oxygen, and 98.5% nitrogen mixture at initial temperatures of 1000, 1100, 1200, and 1500 K.

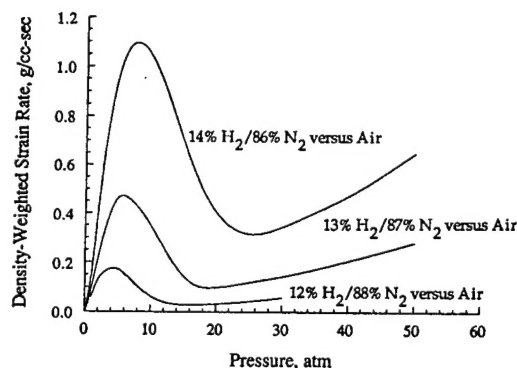


FIG. 6. Air density-weighted extinction strain rates as a function of pressure for 14, 13, and 12% hydrogen-nitrogen mixtures impinging an air stream. Reactant stream exit temperatures are 300 K.

states, while the region to the left represents extinguished states. Thus, an increase in pressure could cause a mixture to change from an extinguished state, to a burning state, and back to an extinguished state.

The above results can best be understood on the basis of the pressure effects on hydrogen oxidation, by examining the maximum hydrogen consumption rate for a *homogeneous* stoichiometric mixture reacting in an adiabatic system *without* mass and heat transport [2]. The maximum reaction rate is defined here as the maximum slope on a hydrogen concentration vs time profile. The computational model describing this closed system at constant pressure is described elsewhere [14]. Figure 5 shows that, for a given initial temperature, the maximum reaction rate, $|dH_2/dt|_{\max}$, first rapidly increases with pressure until a pressure is reached that is indicative of the "extended second limit" [2] shown in Fig. 4. At this pressure, the reaction rate decreases rapidly but then

increases again with pressure. The kinetic details for this behavior are well documented [2]. Specifically, as the pressure is increased at a given temperature, the well-known fast reaction proceeds through the bimolecular branching reactions such as $H + O_2 = OH + O$. The subsequent drop in the maximum reaction rate is the result of the slow reaction that proceeds as a consequence of the chain-inhibiting reaction $H + O_2 + M = HO_2 + M$. It is the competition between the chain branching and inhibiting reactions that explains the second explosion limit and defines the extended second limit. The third explosion limit is the result of the production of H_2O_2 via $2HO_2 = H_2O_2 + O_2$ and $HO_2 + H_2 = H_2O_2 + H$ and the decomposition of H_2O_2 via $H_2O_2 + M = 2OH + M$. Thus, for a given temperature, the reaction rate for pressures above the extended second limit increases again as the reaction path $H + O_2 + M = HO_2 + M$ becomes propagating. The pressure dependency of the maximum reaction rate, as shown in Fig. 5, has been quantitatively confirmed in experimental flow reactor studies of near-adiabatic dilute hydrogen-oxygen mixtures [12]. The behavior of the overall reaction rate with pressure explains the explosion peninsula of the hydrogen-air system of Fig. 4.

For the counterflow configuration studied here, the density-weighted strain rate, ρK , defines the characteristic transport rate. Figure 6 shows the variation of the density-weighted strain rate with pressure for a 13% hydrogen mixture impinging onto an air stream. Similar to the overall reaction rate for the homogeneous hydrogen-oxygen system, the density-weighted strain rate initially increases rapidly with pressure, indicating the increase in the burning intensity. It then decreases at about 7 atm, which is relatively close to the extended second limit at the extinction temperature (1225 K), and eventually increases again. Consequently, for a given density-weighted strain rate, increasing pressure from a low value will bring the system from a state of extinction, to burning, to extinction again, and to burning again.

The final point to note is that, since the above non-monotonic behavior of the explosion limits with pressure is usually associated with low-to-intermediate temperature hydrogen-oxidation kinetics, it may be surprising that it is also observed for burning situations usually associated with high-temperature kinetics. The reason is that, for the highly diluted flames studied here, the flame temperatures are quite low; consequently, a similar change in the rate-controlling steps with pressure responsible for the behavior of the explosion limits is observed. This provides an example where the key reaction steps controlling ignition and extinction events are quite similar.

Summary

In this study, LDV has been employed to experimentally determine the local near-extinction strain

rates for counterflow nonpremixed flames of diluted hydrogen against air at 1.0 and 0.5 atm. A numerical model incorporating a detailed hydrogen-oxidation mechanism was used to describe the experimental results and elucidate the important chemical processes near extinction. The predicted extinction strain rates agree with the experimental values to within 20%. A very important observation has been that the calculated extinction flame temperatures (and strain rates) for given hydrogen concentrations exhibit a nonmonotonic pressure dependence, which is similar to the homogeneous hydrogen-oxygen explosion limits, as well as the calculated ignition temperatures of nonpremixed counterflows of hydrogen-nitrogen and air. The effect of pressure on homogeneous hydrogen-oxidation kinetics effectively explains the nonmonotonic pressure dependency.

Acknowledgments

The authors would like to thank Dr. R. A. Yetter for his valuable insights, Dr. T. G. Kreutz for his assistance, Dr. M. Nishioka for helping generate the extinction turning points shown in Fig. 2, and Mr. J. Sivo for his technical assistance in experimentation. This work was supported by the U.S. Air Force Office of Scientific Research, with an AFRAFT Fellowship awarded to PP.

REFERENCES

1. Yetter, R. A., and Dryer, F. L., *Comb. Sci. Technol.* 79:97-128 (1991).
2. Yetter, R., Dryer, F. L., and Golden, D., *Major Research Topics in Combustion* (M. Y. Hussani, A. Kumar, and R. G. Voight, Eds.) ICASE/NASA Series, Springer-Verlag, New York, 1992, p. 309.
3. Dixon-Lewis, G., and Missaghi, M., *Twenty-Second Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1988, pp. 1461-1470.
4. Gutheil, E., Balakrishnan, G., and Williams, F. A., *Reduced Kinetic Mechanisms for Applications in Combustion Systems*, Lecture Notes in Physics (N. Peters and B. Rogg, Eds.), Springer-Verlag, New York, 1992, p. 177.
5. Gutheil, E., and Williams, F., *Twenty-Third Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1990, pp. 513-521.
6. Darabiha, N., and Candel, S., *Combust. Sci. Technol.* 86:67-85 (1992).
7. Pellett, G. L., Northam, G. B., and Wilson, L. G., "Opposed Jet Diffusion Flames of Nitrogen-Diluted Hydrogen vs Air: Axial LDA and CARS Surveys; Fuel/Air Strain Rates at Extinction," AIAA-89-2522, AIAA/ASME/SAE/ASEE 25th Joint Propulsion Conference, Monterey, CA, July 10-12, 1989.
8. Pellett, G. L., and Northam, G. B., "Strain-Induced Extinction of Hydrogen-Air Counterflow Diffusion

- Flames: Effects of Steam, CO₂, N₂, and O₂ Additives to Air," AIAA-92-0877, 30th Aerospace Sciences Meeting & Exhibit, Reno, NV, January 6, 1992.
9. Kreutz, T. G., and Law, C. K., "Ignition of Nonpremixed Counterflowing Hydrogen versus Heated Air: Computational Study with Detailed Chemistry," *Combust. Flame*, submitted.
 10. Chelliah, H. K., Law, C. K., Ueda, T., Smooke, M. D., and Williams, F. A., *Twenty-Third Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1990, pp. 503-511.
 11. Smooke, M. D., Puri, I. K., and Seshadri, K., *Twenty-First Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1988, pp. 1783-1792.
 12. Kim, T., Master's Thesis, Princeton University, Princeton, NJ (1993).
 13. Kee, R. J., Miller, J. A., and Jefferson, J. H., "CHEMKIN: A General Purpose, Problem Independent, Transportable, Fortran Chemical Kinetic Code Package," Sandia Report, SAND 80-8003, 1980.
 14. Lutz, A. E., Kee, R. J., and Miller, J. A., "SENKIN: A Fortran Program for Predicting Homogeneous Gas Phase Chemical Kinetics with Sensitivity Analysis," SAND Report 87-8248, 1991.
 15. Pirraglia, A. N., Michael, J. V., Sutherland, J. W., and Klemm, R. B., *J. Phys. Chem.* 93:282 (1989).
 16. Sutherland, J. W., Michael, J. V., Pirraglia, A. N., Nesbitt, F. L., and Klemm, R. B., *Twenty-First Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1986, pp. 929-941.
 17. Michael, J. V., and Sutherland, J. W., *J. Phys. Chem.* 92:3853 (1988).
 18. Harding, L. B., and Wagner, A. F., *Twenty-Second Symposium (International) on Combustion*, The Combustion Institute, Pittsburgh, 1988, pp. 983-989.
 19. Slack, M. W., *Combust. Flame* 28:241 (1977).
 20. Tsang, W., and Hampson, R. F., *J. Phys. Chem. Ref. Data* 15:1987 (1986).
 21. Hippler, H., Troe, J., and Willner, J., *J. Chem. Phys.* 93(3):1755 (1990).
 22. Brouwer, L., Cobos, C. J., and Troe, J., *J. Chem. Phys.* 86(11):6171 (1987).
 23. Hippler, H., and Troe, J., *Chem. Phys. Lett.* 192(4):333 (1992).
 24. Warnatz, J., "Survey of Rate Coefficients in C/H/O Systems," in *Combustion Chemistry* (W. C. Gardiner, Jr., Ed.), Springer-Verlag, New York, 1985.